

Figure 8. The HA-IFN α expression cassette in pSAC35. The expression cassette comprises

PRB1 promoter, from S. cerevisiae.

Fusion leader, first 19 amino acids of the HA leader followed by the last 6 amino acids of the MF α -1 leader.

HA-IFNa coding sequence with a double stop codon (TAATAA) ADH1 terminator, from S. cerevisiae. Modified to remove all the coding sequence normaly present in the Hind III/BamHI fragment generally used.

Figure 8

Localisation of 'Loops' based on the HA Crystal Structure which could be used for Mutation/Insertion

1				LQQCPFEDHV HHHHH	
				~ ~	
	I			II	III
51	KTCVADESAE			RETYGEMADC	CAKOEPERNE
	ННННН	ННННН	ННННН	HHHH	H HHHH
					•
101	CELOHKDDND	NT.DRT.VRDEV	DAMCAPERDN	EETFLKKYLY	ELYBBHDAEA
104					
	НННН	·. H	nnnnnnn	нннннннн	nnnn
					•
	*		IV		
151	APELLFFAKR	YKAAFTECCO	AADKAACLLP	KLDELRDEGK	ASSAKORLKC
		нинининн		ннненннннн	
	f1111111111111111111111111111111111111	1111111111111111111	TITITITE	111111111111111111111111111111111111111	***************************************
	,				V
201	ASLQKFGERA				
	ннннн нн	нинининин	нн нн	нинининни	ннинин нн
		,			
		v	· -	VII	
			_		
251		, , , , , , , , , , , , , , , , , , , ,		KPLLEKSHCI	
-	нинининни	ННННН	ННННН	нннннн	H .
301	DIPSTAADEV	ESKDVCKNYA	EAKDVELGME	LYEYARRHPD	YSVVLLLRLA
J U II	НННН	ННННН		НННННН	НННННННН
	nnnn	nnnnnn	nnnnnnn	nnnnnn	nnnnnnn
		VIII			
351	KTYETTLEKC	CAAADPHECY	AKVFDEFKPL	VEEPQNLIKQ	NCELFEQLGE
	ннннннннн			- ннннннннн	

	-		_	4	772
					IX
401	YKFQNALLVR			GKVGSKCC <u>KH</u>	PEAKRMPCAE
•	ннннннннн	нннн н	ННННННННН	ННН	ННННННН
	·				
		. X		ХI	
4 = -	D				
451					A LEVDETYVPK
	нннннннннн	ННННН	нннннннн	ннннннн	ł
					•
501·	EFNAETFTFH	ADICTLSEKE	ROIKKOTALV	ELVKHKPKAT	KEOLKAVMDD
				ННН	
			111111111111111111111111111111111111111	111111	111111111111111111
		XII		· ·	•
551	FAAFVEKCCK	ADDKETCFAE	EGKKLVAASQ	AALGL	
	нининнн	нннн	нинининни	HH	
		********			_
-	. *				
	,		-		
	Loop		Loop		
	I Vals	4-Asn61	VII	Glu280-His	288
	II Thr7	6-Asp89	VIII	Ala362-Glu	
		2-Glu100	IX	Lys439-Pro	
		.70-Ala176			
			X	Val462-Lys	
		47-Glu252	XI	Thr478-Pro	•
	VI- Glu2	266-Glu277	XII	Lys560-Thr	566

Examples of Modifications to Loop IV

a. Randomisation of Loop IV.

IV

151 APELLFFAKR YKAAFTECC<u>Q AADKA</u>CLLP KLDELRDEGK ASSAKQRLKC НИННИНИНИ ИНИНИНИНИ НИНИН НИНИНИНИН НИНИНИНИН

IV

X represents the mutation of the natural amino acid to any other amino acid. One, more or all of the amino acids can be changed in this manner. This figure indicates all the residues have been changed.

b. Insertion (or replacement) of Randomised sequence into Loop IV.



The insertion can be at any point on the loop and the length a length where n would typically be 6, 8, 12, 20 or 25.

Figure 10

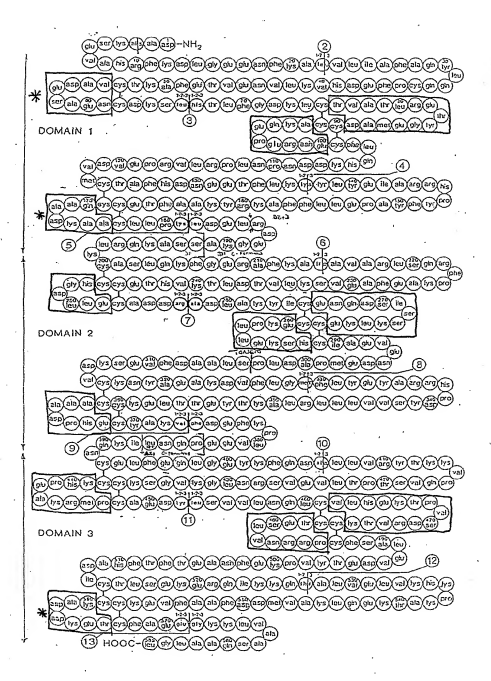
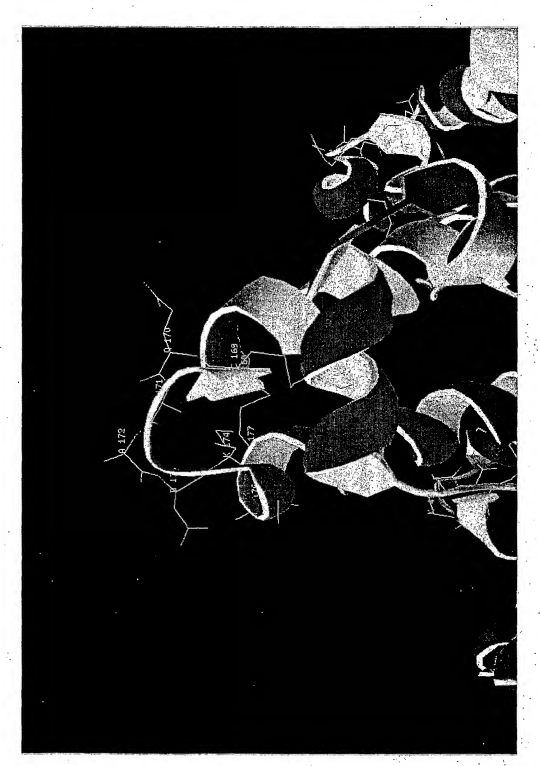


Figure 11



Disulfide bonds shown in yellow

Figure 12: Loop IV Gln170-Ala176

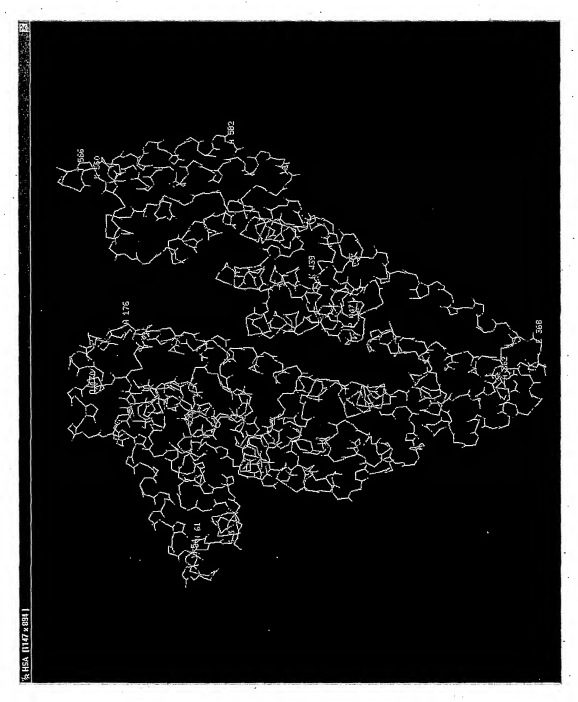


Figure 13: Tertiary Structure of HA

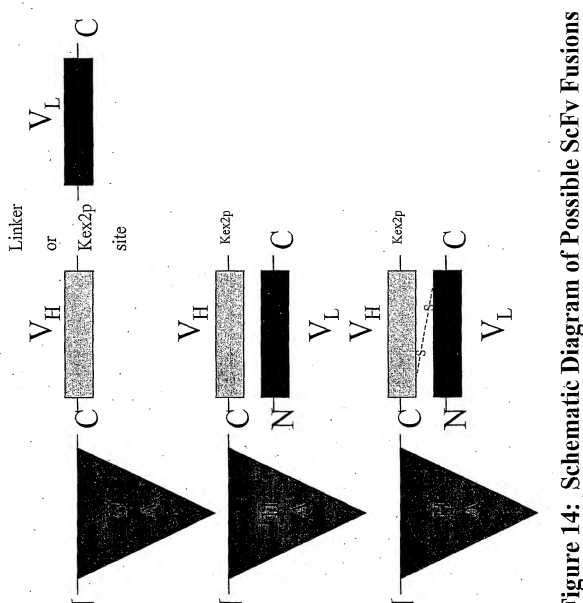


Figure 14: Schematic Diagram of Possible ScFv Fusions (Example is of a C-terminal fusion to HA

			·				
60	120 40	180 60	240	300	360 120	420 140	480 160
AAA K	GTA V	GAA E	. CTT L,	GAA E	GTT V	TAT Y	AĠG R
TTC_{F}	CAT H	GCT A	ACT T	AAT N	GAG	TTA L	AAA K
AAT N	GAT D	TCA S	GCA A	AGA R	CCA	TÀC Y	GCT A
GAA	Gaa E	GAG E	GTT V	GAG E	AGA R	AAA K	· TTT F
GAA E	TTT F	GAT D	ACA T	CCT	$_{\rm V}^{\rm GTG}$	AAA K	TTC
GGA G	CCA	GCT A	TGC D	GAA E	TTG L	TTG L	CTT
TTG	TGT C	GTT V	· TTA L	CAA Q	CGA R	· TTT F	CTC L
GAT D	CAG O	_	AAA K	AAA K	CCC F	ACA T	GAA E
aaa K	CAG Q	ACA T	GAC D	GCA	CTC	GAG E	CCG P
TTT F	CIT	AAA K	GGA G	TGT	AAC N		GCC A
CGG R	TAT Y	GCA A	. TTT 4	TGC	CCA	AAT N	TAT Y
CĄT H	CAG	T'TT F	CTT	3ÁC O	AAC N	GAC D	TTT
GCT A	GCT A	GAA E	ACC	ATG GCT (GAC	CAT H	TAC Y
GTT	TTT	ACT T	CAT H	ATG M	GAT D	TTT F	CCT
GAG E	GCC 	GTA V	CTT	gaa e	AAA K	GCT	CAT H
AGT S	AÌT	GAA E	TCA	GGT	CAC H	ACT T	AGA R
AAG K	TTG	AAT N	AAA K	TAT	CAA Q	TGC C	AGA R
CAC H	GTG V	GTG V	GAC	ACC . T	$\mathop{\text{TTG}}_{L}$	ATG M	GCC A
GCA A	TTG	TTA L	TGT	GAA E	TTC	GTG V	ATT I
1 GAT 1 D	D A	AAA K	AAT	CGT R	TGC C	GAT D	GAA E
н н	7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	121	181	241 81	301	361	421 141
			*			_	

Figure 15A

960 320

180 720 840 280 660 220 780 260 300 GAC D CGC R TGC GCG A GAC AGG GAT GAA E GAG E AAG K GAT D GTG V GAT D AAT TGG W . GAA E GCT GCA AGT S AAA ICC GAA E $_{
m TIC}$ ATC I Ø TCG S ATT AAG K GCT GAA E Ą AGA R GAT D TGC GCA GAT D O ď CAC H GAA E GAG E AAA CAA gaa E GAT D CTCAAG. K AGT S CAC H AGA R CCT P 161 601_. 781 261 661 221

Figure 15B

GAA E 1020 1200 400 CITGGA G TCA S AAA aaa K ${\tt GTG}$ $_{
m LGL}$ CAT CTT L บ AGG $ext{TTT}$ $_{
m LGL}$ Ø ĸ AGA GAG E ACC ACT GAA E ₽ GAT AGC GAA TTC CTT . GĠC GTG GAG E AAG K GTG CTA AAA $^{\mathrm{TGT}}$ ACC T TAT \mathtt{TAT} aaa K AAG သည TAC AAC N GAC CTA GGA ט TAT CAA 闰 Ø AAC TGC ATGAAA K gaa E AGA TCAATG CCC CAT H Щ GIC AAT N CCT P BCG $_{
m GIC}$ GAT D CAG Q AAT N GAG $_{
m GIG}$ GCA GTA CCT P Ø M А AAG GCT A GAG E GCA A GCA GCC A GAA E AAA K ACT T ď 闰 GAG TAC TAC Y \mathbf{TGT} GŢG V CCT 1201 401 361 1141 1261 421 1321 341

Figure 15C

GAC

AGT

500	1560	AAG GCA ACA 1620 K A T 540	1680 560	1740	
CCC AAA P K	GAG E	ACA T	AAG K	CAA Q	
CCC P	AAG K	GCA A	TGC	AGT S	
GTT V	GAG E	AAG K	TGC	GCA A	
ACA TAC T Y	TCT	CCC P	AAG K	GCT A	
ACA T	CTT	AAG K	GAG. E	GTT	1.
GAA	ACA	CAC H	TTT GTA GAG. F V E	CTT	1782
GAT D	TGC C	AAA K	GCA GCT TTT GTA GAG AAG TGC TGC AAG A A F V E K C C K	AAA K	· CAG
GTC V	ATA TGC I C	GTG V	GCT A	aaa. K	TCT
GAA E	GAT D	$_{\rm L}^{\rm CTT}$	GCA	GGT G	GCA
GCT.CTG GAA GTC GAT A L E V D	GCA A	CAA ATC AAG AAA CAA ACT GCA CTT GTG GAA CAC AAG CCC Q I K K Q T A L V E L V K H K P	GCT GTT ATG GAT GAT TTC A V M D D F	GAG GAG GGT AAA AAA CTT GTT GCT GCA AGT CAA E E G K K L V A A S Q	TTA AAA
GCT.	CAT GCA H A	GŢŢ V	GAT	GAG E	TTA
TTT. TCA F S	TTC	$_{\rm L}^{\rm CTT}$	GAT D	GAC GAT AAG GAG ACC TGC TTT GCC D D K E T C F A	CTA CAT
TTT. F	ACC TTC T F	GCA	r ATG M	TTT F	
AGG CGA CCA TGC R R P C	· TTC F	ACT T	GTT V	TGC C	CAT
CCA	aca T	caa o	A GCT C	ACC	TAA
CGA R	GAA	aaa K	GAG CAA CTG AAA E Q L K	GAG E	TTA
AGG R	GCT	AAG K	CTG	AAG K	GGC
GTG AAC V N	AAT N	ATC I	CAA Q	GAT D	TTA
GTG V	TTT F	CAA O	GAG E	GAC D	೨೭೨
${ m TTG}$	GAG E	AGA R	AAA K	GCT	GCT
1441	1501	1561	1621 541	1681 561	1741

Figure 15D

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581 A

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Phe Ala Lys Thr Cys Val Ala Asp Glu Ser Ala Glu Asn Cys Asp Lys 50 55 60

Ser Leu His Thr Leu Phe Gly Asp Lys Leu Cys Thr Val Ala Thr Leu 65 70 75 80

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Arg His Pro Tyr Phe Tyr Ala Pro Glu Leu Leu Phe Phe Ala Lys Arg 145 150 155 160

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Ser Lys Leu Lys Glu Cys Cys Glu Lys Pro Leu Leu Glu Lys Ser His 275 280 285

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Arg	His	Pro	Asp 340	Tyr	Ser	Va1	Val	Leu 345	Leu	Leu	Arg	Leu	Ala 350	Lys	Thr
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Lys 545	Ala	Val	Met	Asp	Asp 550	Phe	Ala	Ala	Phe	Val 555	Glu	Lys	Cys	Cys	Lys 560
Ala	Asp	Asp	Lys	Glu 565	Thr	Cys	Phe	Ala	Glu 570	Glu	Gly	Lys	Lys	Leu 575	Val
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43

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Cys Pro Arg Asn Pro Phe His

INTERNATIONAL SEARCH REPORT

In_____nal application No.
PCT/US01/12009

		
A. CLASSIFICATION OF SUBJECT MATTER IPC(7) :C07K 14/00; C07H 21/04; C12N 15/00; A61K 33 US CL : 435/320.1; 514/2; 530/350; 536/23.4 According to International Patent Classification (IPC) or to be	,	
B. FIELDS SEARCHED		
Minimum documentation searched (classification system follow	ved by classification symbols)	
U.S. : 435/320.1; 514/2; 530/350; 536/23.4		
Documentation searched other than minimum documentation to t	he extent that such documents are included in	n the fields searched
Electronic data base consulted during the international search (name of data base and, where practicable,	search terms used)
MEDLINE, BIOSIS, EMBASE, CAPLUS, various sequen search terms: chimera, fusion, human serum albumin, interf	ce databases	}
C. DOCUMENTS CONSIDERED TO BE RELEVANT		
Category* Citation of document, with indication, where	appropriate, of the relevant passages	Relevant to claim No.
X WO 99/66054 A2 (GENZYME December 1999, see entire document		4,16,17,24,28,30-
Y		37,40-42
		3,15,25,26,29,39 N
X WO 97/24445 A1 (DELTA BIOTEO 1997, seen entire document, especia	CHNOLOGY LIMITED) 10 July ally pp. 3-6, 12, 13, and 19-26	1-4,6,9,10,13- 17,25,28- 34,36,37,39-42
		5,7,8,24,26,2 7,35,38
	•	
X Further documents are listed in the continuation of Box		
Special categories of cited documents: "A" document defining the general state of the art which is not consider	"T" later document published after the in date and not in conflict with the app ed the principle or theory underlying the	olication but cited to understand
to be of particular relevance "B" earlier document published on or after the international filing date	1 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	he claimed invention cannot be ered to involve an inventive step
document which may throw doubts on priority claim(s) or which cited to establish the publication date of another citation or other cited to establish the publication date of another citation or other cited to establish the publication date of another citation or other cited to establish the publication date of another citation or other cited to establish the publication date of another citation or other cited to establish the publication date of another citation or other cited to establish the publication date of another citation or other cited to establish the publication date of another citation or other cited to establish the publication date of another citation or other cited to establish the publication date of another citation or other cited to establish the cited to establish the publication date of another citation or other cited to establish the cited	15	he claimed invention cannot be
special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means	considered to involve an inventiv	e step when the document is ch documents, such combination
"P" document published prior to the international filing date but later the the priority date claimed		
Date of the actual completion of the international search	Date of mailing of the international se	earch report
16 JULY 2001	22 AUG	i 2001
Name, and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT		NY J. DEY
Washington, D.C. 20231	BRONWEN M. LOFARALEGA Telephone No. (703) TECHNOLOG	
Facsimile No. (703) 305-3230	Telephone No. (703) BOSTBISCO	· OFFIRE FILE LOND

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US01/12009

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
Jaivgul y "	Citation of document, their indicators, more appropriate, or an extension	
7	US 5,705,363 A (IMAKAWA, K.) 6 January 1998, see entire document, especially col. 23, line 44-col. 24, line 56.	1,2,4,5-7,10,2 5,33-38,40-42 3,14,15,16,17,24,
		3,14,13,10,17,24,
ζ	US 5,766,883 A (BALLANCE et al) 16 June 1998, see entire document, especially col.1, line 38-54, col. 6, line 53-col 10, line 41, col. 11, line 36-col. 12, line 40.	1,2,4,6,10,13, 14,25,30- 34,36,37,40-42
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(19) World Intellectual Property Organization International Bureau





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- (22) International Filing Date: 12 April 2001 (12.04.2001)
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60/256,931	21 December 2000 (21.12.2000)	US

- (71) Applicants (for all designated States except US): PRINCIPIA PHARMACEUTICAL CORPORATION [US/US]; 2650 Eisenhower Avenue, Building C, Norristown, PA 19403 (US). DELTA BIOTECHNOLOGY LIMITED [GB/GB]; Castle Court, 59 Castle Boulevard, Nottingham NG7 1FD (GB).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): BALLANCE, David, James [GB/US]; 1113 Cymry Drive, Berwyn, PA 19312 (US). SLEEP, Darrell [GB/US]; 66 Ladybay Road, West Bridgford, Nottingham NG2 5DS (GB). TURNER, Andrew, John [GB/US]; Apartment C-28, 305 Conestoga Way, Eagleville, PA 19408 (US). SADEGHI, Homayoun [US/US]; 320 E. Court Street, Doylestown, PA 18901 (US). PRIOR, Christopher, P. [US/US]; 460 Wyldhaven Road, Rosemont, PA 19010 (US).

- (74) Agents: GARRETT, Arthur, S. et al.: Finnegan, Henderson, Farabow, Garrett & Dunner, LLP, 1300 I Street, NW., Washington, DC 20005-3315 (US).
- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW.
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- with (an) indication(s) in relation to deposited biological material furnished under Rule 13bis separately from the description
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28 February 2002

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For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: ALBUMIN FUSION PROTEINS

(57) Abstract: The present invention encompasses albumin fusion proteins. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells. Additionally the present invention encompasses pharmaceutical compositions comprising albumin fusion proteins and methods of treating, preventing, or ameliorating diseases, disorders or conditions using albumin fusion proteins of the invention.

ALBUMIN FUSION PROTEINS

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BACKGROUND OF THE INVENTION

The invention relates generally to Therapeutic proteins (including, but not limited to, a polypeptide, antibody, or peptide, or fragments and variants thereof) fused to albumin or fragments or variants of albumin. The invention further relates to Therapeutic proteins (including, but not limited to, a polypeptide, antibody, or peptide, or fragments and variants thereof) fused to albumin or fragments or variants of albumin, that exhibit extended shelf-life and/or extended or therapeutic activity in solution. These fusion proteins are herein collectively referred to as "albumin fusion proteins of the invention." The invention encompasses therapeutic albumin fusion proteins, compositions, pharmaceutical compositions, formulations and kits. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention using these nucleic acids, vectors, and/or host cells.

The invention is also directed to methods of *in vitro* stabilizing a Therapeutic protein via fusion or conjugation of the Therapeutic protein to albumin or fragments or variants of albumin.

Human serum albumin (HSA, or HA), a protein of 585 amino acids in its mature form (as shown in Figure 15 or in SEQ ID NO:18), is responsible for a significant proportion of the osmotic pressure of serum and also functions as a carrier of endogenous and exogenous ligands. At present, HA for clinical use is produced by extraction from human blood. The production of recombinant HA (rHA) in microorganisms has been disclosed in EP 330 451 and EP 361 991.

The role of albumin as a carrier molecule and its inert nature are desirable properties for use as a carrier and transporter of polypeptides *in vivo*. The use of albumin as a component of an albumin fusion protein as a carrier for various proteins has been suggested in WO 93/15199, WO 93/15200, and EP 413 622. The use of N-terminal fragments of HA

Therapeutic protein may be achieved by genetic manipulation, such that the DNA coding for HA, or a fragment thereof, is joined to the DNA coding for the Therapeutic protein. A suitable host is then transformed or transfected with the fused nucleotide sequences, so arranged on a suitable plasmid as to express a fusion polypeptide. The expression may be effected *in vitro* from, for example, prokaryotic or eukaryotic cells, or *in vivo e.g.* from a transgenic organism.

Therapeutic proteins in their native state or when recombinantly produced, such as interferons and growth hormones, are typically labile molecules exhibiting short shelf-lives, particularly when formulated in aqueous solutions. The instability in these molecules when formulated for administration dictates that many of the molecules must be lyophilized and refrigerated at all times during storage, thereby rendering the molecules difficult to transport and/or store. Storage problems are particularly acute when pharmaceutical formulations must be stored and dispensed outside of the hospital environment. Many protein and peptide drugs also require the addition of high concentrations of other protein such as albumin to reduce or prevent loss of protein due to binding to the container. This is a major concern with respect to proteins such as IFN. For this reason, many Therapeutic proteins are formulated in combination with large proportion of albumin carrier molecule (100-1000 fold excess), though this is an undesirable and expensive feature of the formulation.

Few practical solutions to the storage problems of labile protein molecules have been proposed. Accordingly, there is a need for stabilized, long lasting formulations of proteinaceous therapeutic molecules that are easily dispensed, preferably with a simple formulation requiring minimal post-storage manipulation.

SUMMARY OF THE INVENTION

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The present invention is based, in part, on the discovery that Therapeutic proteins may be stabilized to extend the shelf-life, and/or to retain the Therapeutic protein's activity for extended periods of time in solution, in vitro and/or in vivo, by genetically or chemically fusing or conjugating the Therapeutic protein to albumin or a fragment (portion) or variant of albumin, that is sufficient to stabilize the protein and/or its activity. In addition it has been determined that the use of albumin-fusion proteins or albumin conjugated proteins may reduce the need to formulate protein solutions with large excesses of carrier proteins (such as albumin, unfused) to prevent loss of Therapeutic proteins due to factors such as binding to the container.

The present invention encompasses albumin fusion proteins comprising a Therapeutic protein (e.g., a polypeptide, antibody, or peptide, or fragments and variants thereof) fused to albumin or a fragment (portion) or variant of albumin. The present invention also encompasses albumin fusion proteins comprising a Therapeutic protein (e.g., a polypeptide,

antibody, or peptide, or fragments and variants thereof) fused to albumin or a fragment (portion) or variant of albumin, that is sufficient to prolong the shelf life of the Therapeutic protein, and/or stabilize the Therapeutic protein and/or its activity in solution (or in a pharmaceutical composition) in vitro and/or in vivo. Nucleic acid molecules encoding the albumin fusion proteins of the invention are also encompassed by the invention, as are vectors containing these nucleic acids, host cells transformed with these nucleic acids vectors, and methods of making the albumin fusion proteins of the invention and using these nucleic acids, vectors, and/or host cells.

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The invention also encompasses pharmaceutical formulations comprising an albumin fusion protein of the invention and a pharmaceutically acceptable diluent or carrier. Such formulations may be in a kit or container. Such kit or container may be packaged with instructions pertaining to the extended shelf life of the Therapeutic protein. Such formulations may be used in methods of treating, preventing, ameliotationg or diagnosing a disease or disease symptom in a patient, preferably a mammal, most preferably a human, comprising the step of administering the pharmaceutical formulation to the patient.

In other embodiments, the present invention encompasses methods of preventing treating, or ameliorating a disease or disorder. In preferred embodiments, the present invention encompasses a method of treating a disease or disorder listed in the "Preferred Indication Y" column of Table 1 comprising administering to a patient in which such treatment, prevention or amelioration is desired an albumin fusion protein of the invention that comprises a Therapeutic protein portion corresponding to a Therapeutic protein (or fragment or variant thereof) disclosed in the "Therapeutic Protein X" column of Table 1 (in the same row as the disease or disorder to be treated is listed in the "Preferred Indication Y" column of Table 1) in an amount effective to treat prevent or ameliorate the disease or disorder.

In another embodiment, the invention includes a method of extending the shelf life of a Therapeutic protein (e.g., a polypeptide, antibody, or peptide, or fragments and variants thereof) comprising the step of fusing or conjugating the Therapeutic protein to albumin or a fragment (portion) or variant of albumin, that is sufficient to extend the shelf-life of the Therapeutic protein. In a preferred embodiment, the Therapeutic protein used according to this method is fused to the albumin, or the fragment or variant of albumin. In a most preferred embodiment, the Therapeutic protein used according to this method is fused to albumin, or a fragment or variant of albumin, via recombinant DNA technology or genetic engineering.

In another embodiment, the invention includes a method of stabilizing a Therapeutic protein (e.g., a polypeptide, antibody, or peptide, or fragments and variants thereof) in solution, comprising the step of fusing or conjugating the Therapeutic protein to albumin or a fragment (portion) or variant of albumin, that is sufficient to stabilize the Therapeutic protein.

In a preferred embodiment, the Therapeutic protein used according to this method is fused to the albumin, or the fragment or variant of albumin. In a most preferred embodiment, the Therapeutic protein used according to this method is fused to albumin, or a fragment or variant of albumin, via recombinant DNA technology or genetic engineering.

The present invention further includes transgenic organisms modified to contain the nucleic acid molecules of the invention, preferably modified to express the albumin fusion proteins encoded by the nucleic acid molecules.

BRIEF DESCRIPTION OF THE FIGURES

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Figure 1 depicts the extended shelf-life of an HA fusion protein in terms of the biological activity (Nb2 cell proliferation) of HA-hGH remaining after incubation in cell culture media for up to 5 weeks at 37°C. Under these conditions, hGH has no observed activity by week 2.

Figure 2 depicts the extended shelf-life of an HA fusion protein in terms of the stable biological activity (Nb2 cell proliferation) of HA-hGH remaining after incubation in cell culture media for up to 3 weeks at 4, 37, or 50°C. Data is normalized to the biological activity of hGH at time zero.

Figures 3A and 3B compare the biological activity of HA-hGH with hGH in the Nb2 cell proliferation assay. Figure 3A shows proliferation after 24 hours of incubation with various concentrations of hGH or the albumin fusion protein, and Figure 3B shows proliferation after 48 hours of incubation with various concentrations of hGH or the albumin fusion protein.

Figure 4 shows a map of a plasmid (pPPC0005) that can be used as the base vector into which polynucleotides encoding the Therapeutic proteins (including polypeptides and fragments and variants thereof) may be cloned to form HA-fusions. Plasmid Map key: PRB1p: PRB1 S. cerevisiae promoter; FL: Fusion leader sequence; rHA: cDNA encoding HA: ADH1t: ADH1 S. cerevisiae terminator; T3: T3 sequencing primer site; T7: T7 sequencing primer site; Amp R: β-lactamase gene; ori: origin of replication. Please note that in the provisional applications to which this application claims priority, the plasmid in Figure 4 was labeled pPPC0006, instead of pPPC0005. In addition the drawing of this plasmid did not show certain pertinent restriction sites in this vector. Thus in the present application, the drawing is labeled pPPC0005 and more restriction sites of the same vector are shown.

Figure 5 compares the recovery of vial-stored HA-IFN solutions of various concentrations with a stock solution after 48 or 72 hours of storage.

Figure 6 compares the activity of an HA- α -IFN fusion protein after administration to monkeys via IV or SC.

Figure 7 describes the bioavailability and stability of an HA-α-IFN fusion protein.

Figure 8 is a map of an expression vector for the production of HA- α -IFN.

Figure 9 shows the location of loops in HA.

Figure 10 is an example of the modification of an HA loop.

Figure 11 is a representation of the HA loops.

Figure 12 shows the HA loop IV.

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Figure 13 shows the tertiary structure of HA.

Figure 14 shows an example of a scFv-HA fusion

Figure 15 shows the amino acid sequence of the mature form of human albumin (SEQ ID NO:18) and a polynucleotide encoding it (SEQ ID NO:17).

DETAILED DESCRIPTION

As described above, the present invention is based, in part, on the discovery that a Therapeutic protein (e.g., a polypeptide, antibody, or peptide, or fragments and variants thereof) may be stabilized to extend the shelf-life and/or retain the Therapeutic protein's activity for extended periods of time in solution (or in a pharmaceutical composition) in vitro and/or in vivo, by genetically fusing or chemically conjugating the Therapeutic protein, polypeptide or peptide to all or a portion of albumin sufficient to stabilize the protein and its activity.

The present invention relates generally to albumin fusion proteins and methods of treating, preventing, or ameliorating diseases or disorders. As used herein, "albumin fusion protein" refers to a protein formed by the fusion of at least one molecule of albumin (or a fragment or variant thereof) to at least one molecule of a Therapeutic protein (or fragment or variant of a Therapeutic protein and at least a fragment or variant of human serum albumin, which are associated with one another, preferably by genetic fusion (i.e., the albumin fusion protein is generated by translation of a nucleic acid in which a polynucleotide encoding all or a portion of a Therapeutic protein is joined in-frame with a polynucleotide encoding all or a portion of albumin) or chemical conjugation to one another. The Therapeutic protein and albumin protein, once part of the albumin fusion protein, may be referred to as a "portion", "region" or "moiety" of the albumin fusion protein (e.g., a "Therapeutic protein portion" or an "albumin protein portion").

In one embodiment, the invention provides an albumin fusion protein comprising, or alternatively consisting of, a Therapeutic protein (e.g., as described in Table 1) and a serum albumin protein. In other embodiments, the invention provides an albumin fusion protein comprising, or alternatively consisting of, a biologically active and/or therapeutically active fragment of a Therapeutic protein and a serum albumin protein. In other embodiments, the

invention provides an albumin fusion protein comprising, or alternatively consisting of, a biologically active and/or therapeutically active variant of a Therapeutic protein and a serum albumin protein. In preferred embodiments, the serum albumin protein component of the albumin fusion protein is the mature portion of serum albumin.

In further embodiments, the invention provides an albumin fusion protein comprising, or alternatively consisting of, a Therapeutic protein, and a biologically active and/or therapeutically active fragment of serum albumin. In further embodiments, the invention provides an albumin fusion protein comprising, or alternatively consisting of, a Therapeutic protein and a biologically active and/or therapeutically active variant of serum albumin. In preferred embodiments, the Therapeutic protein portion of the albumin fusion protein is the mature portion of the Therapeutic protein. In a further preferred embodiment, the Therapeutic protein portion of the albumin fusion protein is the extracellular soluble domain of the Therapeutic protein. In an alternative embodiment, the Therapeutic protein portion of the albumin fusion protein is the active form of the Therapeutic protein.

In further embodiments, the invention provides an albumin fusion protein comprising, or alternatively consisting of, a biologically active and/or therapeutically active fragment or variant of a Therapeutic protein and a biologically active and/or therapeutically active fragment or variant of serum albumin. In preferred embodiments, the invention provides an albumin fusion protein comprising, or alternatively consisting of, the mature portion of a Therapeutic protein and the mature portion of serum albumin.

Therapeutic proteins

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As stated above, an albumin fusion protein of the invention comprises at least a fragment or variant of a Therapeutic protein and at least a fragment or variant of human serum albumin, which are associated with one another, preferably by genetic fusion or chemical conjugation.

As used herein, "Therapeutic protein" refers to proteins, polypeptides, antibodies, peptides or fragments or variants thereof, having one or more therapeutic and/or biological activities. Therapeutic proteins encompassed by the invention include but are not limited to, proteins, polypeptides, peptides, antibodies, and biologics. (The terms peptides, proteins, and polypeptides are used interchangeably herein.) It is specifically contemplated that the term "Therapeutic protein" encompasses antibodies and fragments and variants thereof. Thus an albumin fusion protein of the invention may contain at least a fragment or variant of a Therapeutic protein, and/or at least a fragment or variant of an antibody. Additionally, the term "Therapeutic protein" may refer to the endogenous or naturally occurring correlate of a Therapeutic protein.

By a polypeptide displaying a "therapeutic activity" or a protein that is "therapeutically

active" is meant a polypeptide that possesses one or more known biological and/or therapeutic activities associated with a Therapeutic protein such as one or more of the Therapeutic proteins described herein or otherwise known in the art. As a non-limiting example, a "Therapeutic protein" is a protein that is useful to treat, prevent or ameliorate a disease, condition or disorder. As a non-limiting example, a "Therapeutic protein" may be one that binds specifically to a particular cell type (normal (e.g., lymphocytes) or abnormal e.g., (cancer cells)) and therefore may be used to target a compound (drug, or cytotoxic agent) to that cell type specifically.

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In another non-limiting example, a "Therapeutic protein" is a protein that has a biological activity, and in particular, a biological activity that is useful for treating preventing or ameliorating a disease. A non-inclusive list of biological activities that may be possessed by a Therapeutic protein includes, enhancing the immune response, promoting angiogenesis, inhibiting angiogenesis, regulating hematopoietic functions, stimulating nerve growth, enhancing an immune response, inhibiting an immune response, or any one or more of the biological activities described in the "Biological Activities" section below.

As used herein, "therapeutic activity" or "activity" may refer to an activity whose effect is consistent with a desirable therapeutic outcome in humans, or to desired effects in non-human mammals or in other species or organisms. Therapeutic activity may be measured in vivo or in vitro. For example, a desirable effect may be assayed in cell culture. As an example, when hGH is the Therapeutic protein, the effects of hGH on cell proliferation as described in Example 1 may be used as the endpoint for which therapeutic activity is measured. Such in vitro or cell culture assays are commonly available for many Therapeutic proteins as described in the art.

Examples of useful assays for particular Therapeutic proteins include, but are not limited to, GMCSF (Eaves, A.C. and Eaves C.J., Erythropoiesis in culture. In: McCullock EA (edt) Cell culture techniques - Clinics in hematology. WB Saunders, Eastbourne, pp 371-91 (1984); Metcalf, D., International Journal of Cell Cloning 10: 116-25 (1992); Testa, N.G., et al., Assays for hematopoietic growth factors. In: Balkwill FR (edt) Cytokines, A practical Approach, pp 229-44; IRL Press Oxford 1991) EPO (bioassay: Kitamura et al., J. Cell. Physiol. 140 p323 (1989)); Hirudin (platelet aggregation assay: Blood Coagul Fibrinolysis 7(2):259-61 (1996)); IFNα (anti-viral assay: Rubinstein et al., J. Virol. 37(2):755-8 (1981); anti-proliferative assay: Gao Y, et al Mol Cell Biol. 19(11):7305-13 (1999); and bioassay: Czarniecki et al., J. Virol. 49 p490 (1984)); GCSF (bioassay: Shirafuji et al., Exp. Hematol. 17 p116 (1989); proliferation of murine NFS-60 cells (Weinstein et al., Proc Natl Acad Sci 83:5010-4 (1986)); insulin (³H-glucose uptake assay: Steppan et al., Nature 409(6818):307-12 (2001)); hGH (Ba/F3-hGHR proliferation assay: J Clin Endocrinol Metab 85(11):4274-9 (2000); International standard for growth hormone: Horm Res, 51

Suppl 1:7-12 (1999)); factor X (factor X activity assay: Van Wijk et al. Thromb Res 22:681-686 (1981)); factor VII (coagulation assay using prothrombin clotting time: Belaaouaj et al., J. Biol. Chem. 275:27123-8(2000); Diaz-Collier et al., Thromb Haemost 71:339-46 (1994)), or as shown in Table 1 in the "Exemplary Activity Assay" column.

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Therapeutic proteins corresponding to a Therapeutic protein portion of an albumin fusion protein of the invention, such as cell surface and secretory proteins, are often modified by the attachment of one or more oligosaccharide groups. The modification, referred to as glycosylation, can dramatically affect the physical properties of proteins and can be important in protein stability, secretion, and localization. Glycosylation occurs at specific locations along the polypeptide backbone. There are usually two major types of glycosylation: glycosylation characterized by O-linked oligosaccharides, which are attached to serine or threonine residues; and glycosylation characterized by N-linked oligosaccharides; which are attached to asparagine residues in an Asn-X-Ser/Thr sequence, where X can be any amino acid except proline. N-acetylneuramic acid (also known as sialic acid) is usually the terminal residue of both N-linked and 0-linked oligosaccharides. Variables such as protein structure and cell type influence the number and nature of the carbohydrate units within the chains at different glycosylation sites. Glycosylation isomers are also common at the same site within a given cell type.

For example, several types of human interferon are glycosylated. Natural human interferon-α2 is O-glycosylated at threonine 106, and N-glycosylation occurs at asparagine 72 in interferon-a14 (Adolf et al., J. Biochem 276:511 (1991); Nyman TA et al., J. Biochem 329:295 (1998)). The oligosaccharides at asparagine 80 in natural interferon-β1α may play an important factor in the solubility and stability of the protein, but may not be essential for its biological activity. This permits the production of an unglycosylated analog (interferon- $\beta1b$) engineered with sequence modifications to enhance stability (Hosoi et al., J. Interferon Res. 8:375 (1988; Karpusas et al., Cell Mol Life Sci 54:1203 (1998); Knight, J. Interferon Res. 2:421 (1982); Runkel et al., Pharm Res 15:641 (1998); Lin, Dev. Biol. Stand. 96:97 (1998))1. Interferon-γ contains two N-linked oligosaccharide chains at positions 25 and 97, both important for the efficient formation of the bioactive recombinant protein, and having an influence on the pharmacokinetic properties of the protein (Sareneva et al., Eur. J. Biochem 242:191 (1996); Sareneva et al., Biochem J. 303:831 (1994); Sareneva et al., J. Interferon Res. 13:267 (1993)). Mixed O-linked and N-linked glycosylation also occurs, for example in human erythropoietin, N-linked glycosylation occurs at asparagine residues located at positions 24, 38 and 83 while O-linked glycosylation occurs at a serine residue located at

position 126 (Lai et al., J. Biol. Chem. 261:3116 (1986); Broudy et al., Arch. Biochem. Biophys. 265:329 (1988)).

Therapeutic proteins corresponding to a Therapeutic protein portion of an albumin fusion protein of the invention, as well as analogs and variants thereof, may be modified so that glycosylation at one or more sites is altered as a result of manipulation(s) of their nucleic acid sequence, by the host cell in which they are expressed, or due to other conditions of their expression. For example, glycosylation isomers may be produced by abolishing or introducing glycosylation sites, e.g., by substitution or deletion of amino acid residues, such as substitution of glutamine for asparagine, or unglycosylated recombinant proteins may be produced by expressing the proteins in host cells that will not glycosylate them, e.g. in E. coli or glycosylation-deficient yeast. These approaches are described in more detail below and are known in the art.

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Therapeutic proteins corresponding to a Therapeutic protein portion of an albumin fusion protein of the invention include, but are not limited to, plasma proteins. specifically, such Therapeutic proteins include, but are not limited to, immunoglobulins, serum cholinesterase, alpha-1 antitrypsin, aprotinin, coagulation factors in both pre and active forms including but not limited to, von Willebrand factor, fibrinogen, factor II, factor VII, factor VIIA activated factor, factor VIII, factor IX, factor X, factor XIII, c1 inactivator, antithrombin III, thrombin, prothrombin, apo-lipoprotein, c-reactive protein, and protein C. Therapeutic proteins corresponding to a Therapeutic protein portion of an albumin fusion protein of the invention further include, but are not limited to, human growth hormone (hGH), α-interferon, erythropoietin (EPO), granulocyte-colony stimulating factor (GCSF), granulocyte-macrophage colony-stimulating factor (GMCSF), insulin, single chain antibodies, autocrine motility factor, scatter factor, laminin, hirudin, applaggin, monocyte chemotactic protein (MCP/MCAF), macrophage colony-stimulating factor (M-CSF), osteopontin, platelet factor 4, tenascin, vitronectin, in addition to those described in Table 1. These proteins and nucleic acid sequences encoding these proteins are well known and available in public databases such as Chemical Abstracts Services Databases (e.g., the CAS Registry), GenBank, and GenSeq as shown in Table 1.

Additional Therapeutic proteins corresponding to a Therapeutic protein portion of an albumin fusion protein of the invention include, but are not limited to, one or more of the Therapeutic proteins or peptides disclosed in the "Therapeutic Protein X" column of Table 1, or fragment or variable thereof.

Table 1 provides a non-exhaustive list of Therapeutic proteins that correspond to a Therapeutic protein portion of an albumin fusion protein of the invention. The "Therapeutic Protein X" column discloses Therapeutic protein molecules followed by parentheses containing scientific and brand names that comprise, or alternatively consist of, that

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Therapeutic protein molecule or a fragment or variant thereof. "Therapeutic protein X" as used herein may refer either to an individual Therapeutic protein molecule (as defined by the amino acid sequence obtainable from the CAS and Genbank accession numbers), or to the entire group of Therapeutic proteins associated with a given Therapeutic protein molecule disclosed in this column. The "Exemplary Identifier" column provides Chemical Abstracts Services (CAS) Registry Numbers (published by the American Chemical Society) and/or Genbank Accession Numbers ((e.g., Locus ID, NP_XXXXX (Reference Sequence Protein), and XP_XXXXX (Model Protein) identifiers available through the national Center for Biotechnology Information (NCBI) webpage at www.ncbi.nlm.nih.gov) that correspond to entries in the CAS Registry or Genbank database which contain an amino acid sequence of the Therapeutic Protein Molecule or of a fragment or variant of the Therapeutic Protein Molecule. The summary pages associated with each of these CAS and Genbank Accession Numbers are each incorporated by reference in their entireties, particularly with respect to the amino acid sequences described therein. The "PCT/Patent Reference" column provides U.S. Patent numbers, or PCT International Publication Numbers corresponding to patents and/or published patent applications that describe the Therapeutic protein molecule. Each of the patents and/or published patent applications cited in the "PCT/Patent Reference" column are herein incorporated by reference in their entireties. In particular, the amino acid sequences of the specified polypeptide set forth in the sequence listing of each cited "PCT/Patent Reference", the variants of these amino acid sequences (mutations, fragments, etc.) set forth, for example, in the detailed description of each cited "PCT/Patent Reference", the therapeutic indications set forth, for example, in the detailed description of each cited "PCT/Patent Reference", and the activity asssays for the specified polypeptide set forth in the detailed description, and more particularly, the examples of each cited "PCT/Patent Reference" are incorporated herein by reference. The "Biological activity" column describes Biological activities associated with the Therapeutic protein molecule. The "Exemplary Activity Assay" column provides references that describe assays which may be used to test the therapeutic and/or biological activity of a Therapeutic protein or an albumin fusion protein of the invention comprising a Therapeutic protein X portion. Each of the references cited in the "Exemplary Activity Assay" column are herein incorporated by reference in their entireties, particularly with respect to the description of the respective activity assay described in the reference (see Methods section, for example) for assaying the corresponding biological activity set forth in the "Biological Activity" column of Table 1. The "Preferred Indication Y" column describes disease, disorders, and/or conditions that may be treated, prevented, diagnosed, or ameliorated by Therapeutic protein X or an albumin fusion protein of the invention comprising a Therapeutic protein X portion.

					Ductowned Indication V
Ã,	Exemplary	PCT/Patent.	Biological Activity	Exemplary Activity Assay	rreferreu maicauom 1
AS-9-9-0001 P 00		WO8600337 EP103409-A EP155188 US5399684 US5736379 US6025161 US4839283 US4876197-A	Alpha-1-antitrypsin is an an enzyme inhibitor that belongs to the family of serpin serine protease inhibitors. The molecule inhibits the activity of trypsin and elastase.	m m	Emphysema; Infant Respiratory Distress Syndrome; Pulmonary Fibrosis; Respiratory Syncytial Virus Infections; Asthma; Cystic Fibrosis; Genitourinary Disorders; HIV Infections Treatment; Inflammatory Bowel Disorders; Skin Disorders; Viral Hepatitis; Alpha-1 Antitrypsin Deficiency; Adult Respiratory Distress Syndrome
TP 00	LocusID:7450 NP_000543 XP_006947	WO8606096 EP197592 WO9316709-A USS849536 US6008193 US5849702 US5238919	The glycoprotein encoded by this gene H1functions as both an antihemophilic factor carrier and a 1990 20(1):73) platelet-vessel wall mediator in the blood coagulation system. It is crucial to the hemostasis process. Mutations in this gene or deficiencies in this protein result in von Willebrand's disease.	Macroscopic platelet agglutination assay (Wright R. Ann Clin Lab Sci 1990 20(1):73). Collagen binding assay for von Willebrand factor (Favaloro EJ. Thromb Haemost 2000 83(1):127)	Hemophilia; Hemophilia A; von Willebrand disease
7AS-17AS-47AS-47AS-47AS-47AS-47AS-47AS-47AS-4	CAS-155319-91-8 CAS-52014-67-2 LocusID:462 NP_000479 XP_001452	WO9100291 EP568833 GB2116183	Scrpin scrine protease that inhibits thrombin and other proteins involved in blood coagulation	Thrombin activity assay (Verheul et al., Blood 96:4216-4221, 2000)	Sepsis; Thrombosis; Unstable Angina Pectoris; Coagulation disorders; Respiratory Distress Syndrome; Control of blood clotting during coronary artery bypass surgery; Cancer (aaATIII)

Therapeutic Protein X	Exemplary Identifier	PCT/Patent Reference	Biological Activity	Exemplary Activity Assay	Preferred Indication Y
Ano-linoprotein (Ano	CAS-150287-52-8	WO8803166-A	ApoA1 promotes cholesterol	Cholesterol efflux from human	Atherosclerosis; Coronary
E: Apo A4: Apo A1;	LocusID:335	WO9307165-A	efflux from tissues to the liver for	efflux from tissues to the liver for fibroblasts can be directly measured	restenosis;
Apo B)	LocusID:338	US5408038-A.	excretion. ApoA1 is the major	in response to lipid reconstituted	Hyperchotesteroteima,
	LocusID:348	WO9107505-A	protein component of high	ApoAl (J Biol Chem 1996 Oct	Hyperiipidemia; Kaposi's
	NP 000030	W09315198-A	density lipoprotein (HDL) in the	11;271(41):25145-51). The capacity	Sarcoma
	NP_000375	USS472858-A	plasma. ApoA1 is a cofactor for	of ApoB to participate in LDL	
	NP_000032	US5364793-A	lecithin cholesterolacyltransferase	clearance can be evaluated by	
	XP_006435	WO8702062-A	(LCAT), which is responsible for	measurements of ApoB binding to	-
	XP 002288	WO9307165-A	the formation of most plasma	hepatic lipase (J Biol Chem, Vol.	
	XP_008844	WO9856938-A1	cholesteryl esters. Defects in the	273, Issue 32, 20456-20462) and	
	1	US4943527-A	ApoA1 gene are associated with	lipoprotein lipase (J Biol Chem.	
• .			HDL deficiency and Tangier	1995 Apr 7;270(14):8081-6.). Apok	-
	•		disease. ApoB is the main	binding to its receptor can be	
	-	-	apolipoprotein of chylomicrons	measured directly for example as	
•			and low density lipoproteins	illustrated for the liver ApoE	
-		-	(LDL). ApoB binds triglycerides	receptor (J Biol Chem 1986 Mar	*
-		-	and clears LDL from circulation.	25;261(9):4256-67).	
-			ApoE is a component of		
		,	lipoproteins and a ligand for low		
	*		density lipoprotein receptor.		,
			ApoE binds to a specific receptor		• .
			on liver cells and peripheral cells.		
	-		ApoE is essential for the normal		
-		•	linoprotein constituents. Defects	· .	
	•	•			
		-	dysbetalipoproteinemia, or type		
•			III hyperlipoproteinemia (HLP		
		•	III), in which increased plasma		
-		•	cholesterol and triglycerides are		-
			the consequence of impaired		
-	*		clearance of chylomicron and	•	
			V LOCE Politimums:		

Therapeutic Protein X	Exemplary	PCT/Patent	Biological Activity	Exemplary Activity Assay	Preferred Indication Y
-		Reference			
Applagein	CAS-129037-76-9	WO9409036-A	Applaggin (Agkistrodon		Thrombosis; Stroke; Ischemic
(Agkistrodon piscivorus Genbank: A33990	Genbank:A33990	WO9008772-A	piscivorus piscivorus platelet		Heart Disorders
piscivorus (North		W09210575-A	aggregation inhibitor) is a	platelet serotonin release induced by	
American water		JP05255395-A	17,700-Da polypeptide dimer	ADP, gamma-thrombin, and	
moccasin snake):			which a potent inhibitor of	collagen. (Proc Natl Acad Sci U S A	
disulfide-linked Are-			platelet activation. Applaggin	1989 Oct;86(20):8050-4).	
Glv-Asn-containing			blocks platelet aggregation		
dimeric polypeptide)		Ô	induced by ADP, collagen,		•
		•	thrombin, or arachidonic acid.		•
		-	This inhibition is found to		
			correlate with inhibition of		
			thromboxane A2 generation and	•	-
	,		of dense granule release of		
			serotonin.		-
Autocrine motility	LocusID:2821	WO\$707617-A	WO\$707617-A · Glucose phosphate isomerase	Cell motility assay on mouse CT-26 Hemolytic anemia;	Hemolytic anemia;
factor (AMF:	NP 000166	WO9909049-A1	(neuroleukin); neurotrophic factor	(neuroleukin); neurotrophic factor cells (Sun et al., Proc. Natl. Acad. glucosephosphate isomerase	glucosephosphate isomerase
phosphoglucose	XP_012854		and lymphokine; bladder cancer	Sci. USA 96: 5412-5417, 1999; Lin deficiency; Hydrops fetalis	deficiency; Hydrops fetalis
isomerase; glucose			diagnostic.	et al., Mol. Cell. Endocrinol. 84:47-	٠
phosphate isomerase;	-	-		54, 1992)	
neuroleukin)					

ıtion Y	atic ion Injury;								•		•																						
Preferred Indication Y	Angioedema; Pancreatic Disorders; Reperfusion Injury; Transolant Rejection: Vascular	Disorders			•								ā																				
		ar				-							-			•					_		-	-		•					,	-	
Exemplary Activity Assay	C1 inhibitor function can be assessed by measurement of inhibition of complement of	cleavage (J Immunol 1994 Mar	15;152(6):3199-209)		•						•					-					-							-	-				
Biological Activity		 	inhibitor (C1-INH) concentrate	prepared from human plasma is	being successfully used for the	treatment of hereditary	angioneurotic edema. Recently,	C1-INH has been found to be	consumed in severe inflammation	and has been shown to exert	beneficial effects in several	inflammatory conditions such as	human sepsis, post-operative	myocardial dysfunction due to	reperfusion injury, severe	capillary leakage syndrome after	bone marrow transplantation,	reperfusion injury after lung	transplantation, burn, and	cytotoxicity caused by IL-2	therapy in cancer. is a major	inhibitor of two pro-	inflammatory plasma cascade ·	systems, the classical pathway of	complement and the contact	activation system. During the	activation of classical pathway,	C1-INH interacts with the	activated CI and inhibits it.	Interaction of C1-INH with	activated C1 complex leads to the	dissociation of the C1q subunit	and formation
PCT/Patent Reference		3 0	`			•															,	•											
Exemplary Identifier	CAS-80295-38-1 LocusID:710	NP_000053 XP_006339	l				-	-											•									-					
Therapeutic Protein X	1	Berinert; Complement C1 inactivator; C1					•																							•	-	•	

Therapeutic Protein X	Exemplary Identifier	PCT/Patent Reference	Biological Activity	Exemplary Activity Assay	Preferred Indication Y
Coagulant Complex (Anti-Inhibitor Coagulant Complex; FEIBA VH;					Control of spontaneous bleeding in hemophilia A and B; prevention of bleeding in patients on Factor VIII inhibitors
C-reactive protein	LocusID: 1401 NP_000558 XP_001859	WO9201364-A WO9505394-A	Acute-phase serum protein that binds microbial polysaccharides and ligands on damaged cells, activates the classical complement pathway	Platelet activation (Simpson RM. Immunology 1982 47(1):193) Platelet aggregation (Cheryk LA.Vet Immunol Immunopathol 1996 52:27). Increased production of IL-1 alpha, IL-1 beta, and TNF-alpha in macrophages (Galve-de Rochemonteix B. J Leukoc Biol 1993 53:439)	
EPO (Erythropoietin; Epoetin alfa; Epoetin beta; Gene-activated erythropoietin; Darbepoetin-alpha; NESP; Epogen; Procrit; Eprex; Erypo; Espo; Epoimmun; EPOGIN; NEORECORMON; HEMOLINK; Dynepo; ARANESP)	CAS-113427-24-0 CAS-122312-54-3 LocusID:2056 NP 000790 XP 011627	WO9902710-A1 WO8502610-A WO8603520-A WO9206116- A WO9206116-A US5985607-A EP232034	Hormone that senses and regulates the level of oxygen in the blood by modulating the number of circulating erythrocytes	Cell proliferation assay using a erythroleukemic cell line TF-1. (Kitamura et al. 1989 J.Cell. Physiol. 140:323)	Anemia; Bleeding Disorders

tition CAS- Locus c; NP_0 XP_0		Reference			
Factor IX (Coagulation CAS-1810 factor IX (human); LocusID:2 Factor IX Complex; NP_0001 Christmas factor; XP_0102 plasma thromboplastin					
factor IX (human); LocusID:2 Factor IX Complex; NP_00015 Christmas factor; XP_0102 plasma thromboplastin	2158	WO8505125-A		Factor IX clotting activity: Valder	Hemophilia B; bleeding;
	2170	WO8505376-A	hat	R. et al., 2001 "Posttranslational	Factor LX deliciency;
	24	W09747737-A1		modifications of recombinant	Christmas disease; bieeding
	70	EP162782-A		myotube-synthesized human factor	episodes in patients with
1 4 (•	WO8400560-A	converted to an active form by	IX" Blood 97: 130-138.	Tactor VIII Infilipitor of Factor
component(PTC);		US4994371-A	factor XIa, which excises the		VII deficiency
prothrombin complex			activation peptide and thus		,
concentrate (PCC):		.	generates a heavy chain and a		
Nonacog alpha.			light chain held together by one	•	
MONONINE:			or more disulfide bonds. In the		
AT PHANINE-SD:		•	blood coagulation cascade,		
BEBULIN: PROPLEX-	-	-	activated factor IX activates factor		
T: KONYNE:	8		X to its active form through		
PROFIL NINE SD:			interactions with Ca+2 ions,		
BeneFIX: IMMUNINE			membrane phospholipids, and		3
(MA)			factor VIII. Alterations of this		
			gene, including point mutations,		
•			insertions and deletions, cause		
	•	,	factor IX deficiency, which is a		
			recessive X-linked disorder, also		
	•	4	called hemophilia B or Christmas		
			disease.		

PCT Ref	l/Patent ference	Biological Activity	Exemplary Activity Assay	Preferred Indication Y
WO8400560-A	S60-A	Coagulation factor VII is a	Coagulation Assay using	Bleeding Disorders; Coronary
WO9323074-A	74-A	vitamin K-dependent factor	Prothrombin Clotting Time	Restenosis; Hemophilia A and
JS5997864-A	4-A	essential for hemostasis. This	(Belaaouaj AA et al., J. Biol. Chem. B; Liver Disorders;	B; Liver Disorders;
JS5580560-A	P-4	factor circulates in the blood in a	275: 27123-8, 2000, Diaz-Collier JA Thrombosis, Vascular	I hrombosis; Vascular
JS4994371-A	Ą.	zymogen form, and is converted	et al., Thromb Haemost 71: 339-46; Restenosis; Surgery-related	Kestenosis; Surgery-related
EP200421-A		to an active form by either factor	1994).	hemorrhagic episodes
W09427631-A	Y-	IXa, factor Xa, factor XIIa, or		
WO9309804-A	Y -	thrombin by minor proteolysis.	•	
		Upon activation of the factor VII,		
		a heavy chain containing a		
		catalytic domain and a light chain		
	•	containing 2 EGF-like domains		
	•	are generated, and two chains are		
		held together by a disulfide bond.		
		In the presence of factor III and	•	
		calcium ions, the activated factor		
		then further activates the		
		coagulation cascade by converting		
		factor IX to factor IXa and/or		•
		factor X to factor Xa. Defects in		
		this gene can cause coagulopathy.		

	T		ပ္													-						-						
Preferred Indication Y	·	Hemophilia A; Hemophilia;	Surgery-related hemorrhagic	episodes	ı				•			*			;	-												
Exemplary Activity Assay		Development of a simple	chromogenic factor VIII assay for	clinical use.	Wagenvoord RJ, Hendrix HH,	Hemker HC. Haemostasis	1989;19(4):196-204.	•							-			-						•				
Biological Activity	•	This gene encodes coagulation	factor VIII, which participates in	the intrinsic pathway of blood	coagulation; factor VIII is a	cofactor for factor IXa which, in	the presence of Ca+2 and	phospholipids, converts factor X	to the activated form Xa. This	gene produces two alternatively	spliced transcripts. Transcript	variant 1 encodes a large	glycoprotein, isoform a, which	circulates in plasma and	associates with von Willebrand	factor in a noncovalent complex.	This protein undergoes multiple	cleavage events. Transcript	variant 2 encodes a putative small	protein, isoform b, which	consists primarily of the	phospholipid binding domain of	factor VIIIc. This binding domain	is essential for coagulant activity.	Defects in this gene results in	hemophilia A, a common	recessive X-linked coagulation	disorder
PCT/Patent	Reference	WO9621035-A2	WO9703195-A1	WO9800542-A2	EP160457-A	EP160457-A	WO9959622-A1	EP253455-A				•		-												·.		,
Exemplary		CAS-139076-62-3	LocusID:2157	NP 000123	XP 013124	I		5						•	-						-							
Therapeutic Protein X		Factor VIII (Factor VIII; CAS-139076-62-3	Octocog alfa;	ılfa;		lic factor:	Nordiate: ReFacto:	Kogenate: Kogenate	SF. Helixate:	Recombinate)							-											

Exemplary	PCT/Patent	Biological Activity	Exemplary Activity Assay	Preferred Indication Y
Identifier	Reference			
l	WO9204378-A	Encodes the vitamin K-dependent	Encodes the vitamin K-dependent FACTOR X ACTIVITY ASSAY.	Factor X deficiency; Stuart-
•	WO9309804-A	coagulation factor X precursor of	coagulation factor X precursor of Vall Wijk Eivi et al. A laptu manuar i rower factor centrefer.	hemorrhage; menorrhagia;
•	11111111111	This factor precursor is converted		hematuria; hemarthrosis
		to a mature two-chain form by		
		the excision of the tripeptide		
		RKR. Two chains of the factor		
		are held together by 1 or more	• ,	
		disulfide bonds; the light chain	•	
	ä	contains 2 EGF-like domains,		
		while the heavy chain contains		
		the catalytic domain which is		
		structurally homologous to those		
	-	of the other hemostatic serine	•	
		proteases. The mature factor is		,
	-	activated by the cleavage of the	,	
		activation peptide by factor IXa		
	•	(in the intrinsic pathway), or by		
	•	factor VIIa (in the extrinsic		
	•	pathway). The activated factor		,
		then converts prothrombin to		•
		thrombin in the presence of factor		
		Va, Ca+2, and phospholipid		
		during blood clotting. Mutations		
	•	of this gene result in factor X		
		deficiency, a hemorrhagic	•	
		condition of variable severity.		

Identifier LocusID:2162 LocusID:2163 LocusID:2164 LocusID:2165 NP 000120 NP 001985 XP 004467 XP 001350	1er 5	Reference			100
	5 4 3				
	£ 4 %	W09116931-A	Coagulation factor XIII is the last BLOOD COAGULATION ASSAY.	BLOOD COAGULATION ASSAY.	Factor XIII deficiency;
LocusID:21 LocusID:21 NP_000120 NP_001985 XP_004467 XP_001350	4.5	EP494702-A	zymogen to become activated in	Karpati L, Penke B, Katona E,	bleeding tendency; detective
LocusID:21 NP_000120 NP_001985 XP_004467 XP_001350	\$	US7425887-A		Balogh I, Vamosi G, Muszbek L.	wound healing; habitual
NP_000120 NP_001985 XP_004467 XP_001350	•	WO9102536-A		A modified, optimized kinetic	abortion
NP_001985 XP_004467 XP_001350		WO9918200-A	_	photometric assay for the	
XP_004467 XP_001350				determination of blood coagulation	
XP_001350			A subunits have catalytic	factor XIII activity in plasma. Clin	
			function, and the B subunits do	Chem. 2000 Dec;46(12):1946-55.	
			not have enzymatic activity and		,
		•	may serve as a plasma carrier		•
•			molecules. Platelet factor XIII is	,	
-			comprised only of 2 A subunits,		
			which are identical to those of	-	•
•			plasma origin. Upon activation		
			by the cleavage of the activation		
		1	peptide by thrombin and in the		
		-	presence of calcium ion, the		
			plasma factor XIII dissociates its		-
			B subunits and yields the same		
	•		active enzyme, factor XIIIa, as		
•	-	j	platelet factor XIII. This enzyme	•	
and the same of th			acts as a transglutaminase to		
· · ·	- •	•	catalyze the formation of gamma-		
•			glutamyl-epsilon-lysine		
			crosslinking between fibrin	٠	
			molecules, thus stabilizing the		
			fibrin clot. Factor XIII deficiency		
			is classified into two categories:	,	
	*		type I deficiency, characterized by	,	
			the lack of both the A and B	,	
* * * * * * * * * * * * * * * * * * * *	-		subunits; and type II deficiency,		
			characterized by the lack of the A	;	•
•			subunit alone		

	•	
Preferred Indication Y	Tissue adhesion; thrombosis; bleeding disorders; wounds; thrombocytopenia; dysfibrinogenemia; hypofibrinogenemia; renal amyloidosis; thrombosis; dysprothrombinemia	
Exemplary Activity Assay	Fibrinogen assay: Halbmayer WM, Haushofer A, Schon R, Radek J, Fischer M. Comparison of a new automated kinetically determined fibrinogen assay with the 3 most used fibrinogen assays (functional, derived and nephelometric) in Austrian laboratories in several clinical populations and healthy controls. Haemostasis 1995 May-Jun;25(3):114-23. Tan V, Doyle CJ, Budzynski AZ. Comparison of the kinetic fibrinogen assay with the von Clauss method and the clot recovery	method in plasma of patients with conditions affecting fibrinogen coagulability. Am J Clin Pathol. 1995 Oct;104(4):455-62. Lawrie AS, McDonald SJ, Purdy G, Mackie IJ, Machin SJ. Prothrombin time derived fibrinogen determination on Sysmex CA-6000. J Clin Pathol. 1998 Jun;51(6):462-6. Aprotinin assay: Cardigan RA, Mackie IJ, Gippner-Steppert C, Jochum M, Royston D, Gallimore MJ. Determination of plasma aprotinin levels by functional and immunologic assays. Blood Coagul Fibrinolysis. 2001 Jan;12(1):37-42. Thrombin assay: Syed S, R[4]C PD, Kulczycky M, Sheffield WP. Potent antithrombin activity and delayed clearance from the circulation characterize recombinant hirudin genetically fused to albumin. Blood. 1997 May 1;89(9):3243-52.
Biological Activity	Following vascular injury, fibrinogen assay: Halbmayer W fibrinogen is cleaved by thrombin to form fibrin, which is the most abundant component of blood clots. In addition, various and fibrin regulate cell adhesion and fibrin regulate cell adhesion and spreading, display activities, and are mitogens for several cell types. Coagulation of the coagulation cascade, which clots to form thrombin in the first step of the coagulation cascade, which clots abundant to form thrombin in the first step of the coagulation cascade, which	ultimately results in the stemming of blood loss. F2 also plays a role in maintaining vascular integrity.
PCT/Patent	US6083902-A WO9523868-A1 WO9523868-A1 WO9523868-A1 WO952968-A1 US6083902-A WO9523868-A1 WO9523868-A1 WO9528946-A1 WO9528946-A1 US502034-A US502034-A US502034-A	
Exemplary	LocusID:2243 LocusID:2244 LocusID:2246 LocusID:2266 LocusID:2147 NP_000499 NP_068657 NP_005132 NP_00500 NP_068656 NP_006497	
Therapeutic Protein X	Fibrinogen; thrombin; aprotinin (Human fibrinogen; human thrombin; aprotinin; and calcium chloride; synthocytes; FAMs; BERIPLAST-P)	

Preferred Indication Y		Bone Marrow Disorders, Bone marrow transplant; Chemoprotection; Hepatitis C; HIV Infections; Lung Cancer; Malignant melanoma; Mycobacterium avium complex; Mycoses; Myeloid Leukemia; Neonatal infections; Neutropenia; Oral mucositis; Prostate Cancer; Stem Cell Mobilization; Vaccine Adjuvant; Venous Stasis Ulcers; Prevention of neutropenia; Acute myelogenous leukemia; Hematopoietic progenitor cell mobilization; Non-Hodgkin's lymphoma; Acute lymphoblastic leukemia; Hodgkin's disease; Accelerated myeloid recovery; Xenotransplant rejection
Exemplary Activity Assay	Proliferation of murine NFS-60 cells (Weinstein et al, Proc Natl Acad Sci U S A 1986; 83, pp5010-4)	Colony Stimulating Assay: Testa, N.G., et al., "Assays for hematopoietic growth factors." Balkwill FR (edt) Cytokines, A practical Approach, pp 229-44; IRL Press Oxford 1991.
Biological Activity	Stimulates the proliferation and differentiation of the progenitor cells for granulocytes	Regulates hematopoietic cell differentiation, gene expression, growth
PCT/Patent Reference	WO8604506-A EP220520-A WO8604506-A WO8701132-A US6054294-A	WO8805786 WO86003225 US5391706 US5545536
Exemplary	CAS-121181-53-1 CAS-135968-09-1 CAS-130120-55-7 CAS-130120-54-6 CAS-134088-74-7 LocusID:1440 NP_000750 XP_008227	CAS-99283-10-0 CAS-123774-72-1 CAS-60154-12-3 CAS-137463-76-4 LocusID:1437 NP_000749 XP_003751
Therapeutic Protein X	G-CSF (Granulocyte colony-stimulating factor; Granulokine; KRN 8601; Filgrastim; Lenograstim; Meograstim; Nartograstim; Nartograstim; Oran; GRANOCYTE; Granulokine; Neutrogin; Neu-up; Neutrogin; Neu-up;	GM-CSF (Granulocyte-macrophage colony-stimulating factor; rhuGM-CSF; BI 61012; Prokine; Molgramostim; Sargramostim; GM-CSF/IL 3 fusion; Milodistim; Leucotropin; PROKINE; LEUKOMAX; Interberin; Leukine; Leukine Liquid; Pixykine)

Therapeutic Protein X	Exemplary	PCT/Patent Reference	. Biological Activity	Exemplary Activity Assay	Preferred Indication Y
	Identinei	and latest			Dynosius to Usnotitic B
II andtitica IC (II cantitio		•			Exposure to trepatitis D
Hepannis IO (nepannis	•		•		(HRsAp) or Henatitis C:
B Immine Globiilin:					(2.13)
Turning Clocking			•		perinatal exposure of infants
Hepatitis C immune					with HRV or HCV infected
alohulin. IICVIG.					ואומו זודה ל סו ווכל מווימנים
giodulii, 110 v.C.					mothers;; sexual or household
BAYHEP; NABI-HB;	•	•	-		exposure to netient with soute
(dro v vio ra v id					exposure to patient with acute
NADI-CIVACIN					HBV or HCV

Preferred Indication Y	Alopecia; Cancer, Chemoprotection; Cirrhosis; Haematological disorders;	otection																					-	•	-		
	er	unal of Cell Radioprotection		chanism of chanism of change in the change of	uppl. 59: 63-		erization of	3	iochemical	1); Dowrick	ffects major	etal	al cells	991), parison of	chemical	scatter factor	actor are	nal of Cell	91); Gherardi	catter factor,	ic protein that	eractions and	s of the	cience (USA)	<u>-</u>		
Exemplary Activity Assay	Adams JC et al Production of scatter factor by ndk, a strain of epithelial cells, and inhibition of scatter factor	activity by suramin: Journal of Cell Science 98: 385-94 (1991); Bhargava	MM et al Purification,	characterization and mechanism of action of scatter factor from human	placenta. Experientia Suppl. 59: 63-	75 (1991); Coffer A et al	Purification and characterization of	ras-transformed NIH-3T3	conditioned medium. Biochemical	Journal 278: 35-41 (1991); Dowrick	PG et al Scatter factor affects major	changes in the cytoskeletal	organization of epitheli	Cytokille 3. 239-310 (1331), Furlong R A et al Comparison of	biological and immunochemical	properties indicates that scatter factor	and hepatocyte growth factor are	indistinguishable. Journal of Cell	Science 100: 173-7 (1991); Gherardi	E et al Purification of scatter factor,	a fibroblast-derived basic protein that	modulates epithelial interactions and	movement. Proceedings of the	National Academy of S			
Biological Activity	HGF disrupts desmosomal junctions between epithelial cells and induces a motile fibroblast-	like phenotype in individual cells. The factor therefore also	influences the invasive growth of	tumor cells derived from	involved also in processes of	Wound healing and early	embryonic development. For	some cell types including keratinocytes and mammary	epithelial cells HGF is merely a	motility factor. It is also an	autocrine modulator that	influences the motility of the		mitogen for nepatocytes and also	hepatocyte growth factor). HGF	binds to heparin and this may be	important for its activities in	vivo. The actions of HGF are	inhibited by Suramin . HGF has	been shown to be an	Angiogenesis factor in vivo. It	induces cultured microvascular	endothelial cells to accumulate	and secrete significantly increased inational Academy of Science (USA)	quantities of urokinase, an enzyme associated with	development of an invasive	endothelial phenotype during
PCT/Patent Reference	JP03130091-A JP10070990-A WO9323541-A		- %		3						•											,	-	•	•	,	
Exemplary Identifier	LocusID:3082 Genbank:CAA34387			•		•	•		•		•	•	-													ě	
Therapeutic Protein X	Hepatocyte growth factor (HGF, Scatter factor: SF: HGF/SF)				8.										•				•								•

y Preferred Indication Y	ed Coronary restenosis; Deep Say Vein Thrombosis; Disseminated Intravascular Coagulation; Heparin-induced thrombocytopenia and thrombosis syndrome; Myocardial infarction; Unstable Angina Pectoris; Anticoagulant in adults suffering from acute coronary syndrome; Thrombosis; Veinous Thrombosis
Exemplary Activity Assay	Hirudin activity can be measured using a platelet aggregation assay (Blood Coagul Fibrinolysis 1996 Mar,7(2):259-61).
Biological Activity	Hirudin is a potent anticoagulant Hirudin activity can be measured using a platelet aggregation assay (Blood Coagul Fibrinolysis 1996 Mar,7(2):259-61).
PCT/Patent Reference	WO8504418-A EP200655-A EP503829-A WO9201712-A WO9201712-A EP340170-A EP341215-A WO9207874-A WO9201712-A
Exemplary	CAS-138068-37-8 W08504418-A CAS-120993-53-5 EP200655-A CAS-8001-27-2 EP503829-A Genbank: AAA29195 W09201712-A Genbank: AAA01384 W09201712-A EP340170-A EP341215-A W09201712-A W09201712-A
Therapeutic Protein X	Hirudin (Lepirudin; Desirudin; Refludan; Revasc)

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Preferred Indication Y	Acromegaly; Growth failure; Growth failure and endogenous growth hormone replacement; Growth hormone deficiency; Growth failure and growth retardation Prader- Willi syndrome in children 2 years or older; Growth deficiencies; Postmenopausal osteoporosis; burns; cachexia; cancer cachexia; dwarfism; metabolic disorders; obesity; renal failure; Turner's Syndrome; fibromyalgia; fracture treatment; frailty	
ferred In	Acromegaly; Growth fail Growth fail Growth failure and endogenous growth horn replacement, Growth horn deficiency; Growth failun growth retardation Pradel Willi syndrome in childtyears or older; Growth deficiencies; Postmenops osteoporosis; burns; cach cancer cachexia; dwarfism metabolic disorders; obe renal failure; Turner's Syndrome; fibromyalgia fracture treatment; frailty	* : ' '
Pre		. 0
Assay	Ba/F3-hGHR proliferation assay, a novel specific bioassay for serum human growth hormone. J Clin Endocrinol Metab 2000 Nov;85(11):4274-9 Plasma growth hormone (GH) immunoassay and tibial bioassay, Appl Physiol 2000 Dec;89(6):2174-8 Growth hormone (hGH) receptor mediated cell mediated proliferation, Growth Horm IGF Res 2000 Oct;10(5):248-55 International standard for growth hormone, Horm Res 1999;51 Suppl	
Exemplary Activity Assay	Ba/F3-hGHR proliferation assay, a novel specific bioassay for serum human growth hormone. J Clin Endocrinol Metab 2000 Nov;85(11):4274-9 Plasma growth hormone (GH) immunoassay and tibial bioassay, Appl Physiol 2000 Dec;89(6):2174 8 Growth hormone (hGH) receptor mediated cell mediated proliferatio Growth Horm IGF Res 2000 Oct;10(5):248-55 International standard for growth hormone, Horm Res 1999;51 Suppl 1:7-12	
mplary	Ba/F3-hGHR proliferatiovel specific bioassay human growth hormone Endocrinol Metab 2000 Nov;85(11):4274-9 Plasma growth hormone immunoassay and tibia Appl Physiol 2000 Dec 8 Growth hormone (hGH) mediated cell mediated Growth Horm IGF Res Oct;10(5):248-55 International standard fi hormone, Horm Res 19 1:7-12	
Exe	Ba/F3-h novel sp human E Endocrii Nov;85(Plasma i immuno Appl Ph 8 Growth mediate Growth Oct;10(:	-
ty	Plays an important role in growth Ba/F3-hGHR proliferation assay, a control; binds 2 GHR molecules human growth hormone. J Clin human growth hormone. J Clin Endocrinol Metab 2000 Nov;85(11):4274-9 Plasma growth hormone (GH) immunoassay and tibial bioassay, Appl Physiol 2000 Dec;89(6):2174 8 Growth hormone (hGH) receptor mediated cell mediated proliferation Growth Horm IGF Res 2000 Oct;10(5):248-55 International standard for growth hormone, Horm Res 1999;51 Suppl 1:7-12	
Biological Activity	ant role i GHR m nal transc r dimeriz	
Biologic	Plays an important role in growt control; binds 2 GHR molecules and induces signal transduction through receptor dimerization	· .
	Plays a contro and inc through	
PCT/Patent Reference	WO9418227-A WO9005185-A WO9520398-A EP245138-A WO8605804-A WO90418227-A1 US6194176-A US6194176-A US6194176-A US6194176-A US61977089 US580723 US580723 US580723 US5955346 US6013478	WO9004788-A WO9418227-A1 US6110707-A US4977089 US5580723 US558346 US6013478 WO8605804 US6110707 US5580723 US5580723 US6113478 WO8605804 WO8605804 WO9004788-A WO9004788-A US6110707-A US6110707-A US6110707-A US6110707-A US6110707-A US6580723
PCT Red	WO9418227- WO9005185- WO9520398- EP245138-A- WO8605804- WO9004788- WO9018227- US61194176- WO86013579- US6110707- US6110707- US6110707- US5580723- US5580723- US5580723- US5580723- US5580723- US5580723- US613478- WO8605804-	WO9004788 WO9418227 US6110707- US4977089 US5580723 US5955346 US6013478 WO8605804 US5580723 US5955346 US6013478 WO9004788 WO9014788 WO9418227 US6110707- US6110707- US6110707- US6110707- US6110707-
lary	-87-3 -01-5 88 89 ,	
Exemplary Identifier	CAS-82030-87-3 CAS-12629-01-5 LocusID:2689 NP 000506 NP 072053 NP 072054 NP 072056 NP 072056 NP 072050 NP 072050 NP 072050 NP 072050 NP 072050	
n X	25.5	
ic Protei	wth egyisam somatrol i. Somatrol i. N.; BIO-DE; A; AQ; IN; OPIN; PIN; PIN; EROSTI	
Therapeutic Protein X	Human growth hormone (Pegvisamont; Somatropin; TROVERT; PROTROPIN; BIO-TROPIN; HUMATROPIN; NUTROPIN; NUTROPIN; NORDITROPIN; SAIZEN; SEROSTIM)	
E	# 5 S F E F E E E E E E E	

Therapeutic Protein X	Exemplary	PCT/Patent	Biological Activity	Exemplary Activity Assay	Preferred Indication Y
	Identifier	Reference		-	- 3
Insulin (Human insulin: ICAS-11061-68-0	CAS-11061-68-0	WO200040613-	Insulin is a heterodimeric	Insulin activity may be assayed in	Hyperglycemia; Diabetes
Insulin aspart Insulin CAS-116094-23-6	CAS-116094-23-6	Al	polypeptide hormone involved in	polypeptide hormone involved in vitro using a [3-H]-glucose uptake	mellitus; Type I diabetes and
Glaroine: Insulin lishro: CAS-133107-64-9	CAS-133107-64-9	EP37723-A	carbohydrate metabolism. After	assay. (J Biol Chem 1999 Oct 22;	type 2 diabetes
1 vs-R28 Pro-R29:	CAS-160337-95-1	EP55942-A	removal of the precursor signal	[274(43):30864-30873).	
	LocusID:3630	US4431740-A	peptide, proinsulin is post-		
diarginylinsulin; Des-	NP 000198	US4430266-A	translationally cleaved into two		
B26-B30-insulin-B25-	XP_006400	US4624926-A	chains (peptide A and peptide B)		,
amide: Insulin detemir;	1	US5077204-A	that are covalently linked via two		
LABI; NOVOLIN;	,	US5840542-A	disulfide bonds. Binding of this		-
NOVORAPID;		US6110707-A	mature form of insulin to the		
HUMULIN;		WO9200322-A	(insulin receptor (INSR)		
NOVOMIX 30;	-	•	stimulates glucose uptake.		
VELOSULIN;					
NOVOLOG; LANTUS;					
(liletin; humalog;			•		
MACRULIN;	•		•	•	2
EXUBRA; INSUMAN;	•	-			
ORALIN; ORALGEN;	-	•			
HUMAHALE;	•				
HUMAHALIN)	•				

Preferred Indication Y	Immune deficiencies;	agammaglobulinemia;	inypogaminagioominemia; imminodeficient states and	bacterial infections; Kawasaki	Syndrome; Hepatitis A;	measles varicella; rubella;	immunoglobulin deficiency;	idiopathic thrombocytopenic	purpura; primary humoral	immunodeficiency states;	bone marrow transplantation;	pediatric HIV infection;	Guillain-Barre syndrome;	emonic initatimatory	demyetmating polyneuropathy: multifocal	neuropathy: dermatomyositis:	amyotrophic lateral sclerosis;	inclusion-body myositis;	Lambert-Eaton myasthenic	syndrome; Rasmussen	syndrome; West syndrome;	intractable childhood epilepsy;	Lennox-Gastaut syndrome;	polymyositis; relapsing-	remitting multiple sclerosis;	opue neums, sunt-man	cerebellar degeneration:	paraneoplastic	encephalomyelitis and sensory	neuropathy; systemic	vasculitis; myelopathy	loo il nomina attita patoroccori
Exemplary Activity Assay								•									٠													,	•	
Biological Activity	Regulates hematopoietic cell	differentiation, gene expression,	growth					•							,	-					2		•						-			
PCT/Patent Reference						•											*		•	-		•			-							
Exemplary Identifier			•											•							•	•		,		-		•			,	
Therapeutic Protein X	IVIG (Intravenous	Immune Globulin;	VENOGLOBULIN-S;	PANGLOBOLIN,	GAMMAR-P	GAMMAGARD S/D;	IVEEGAM; BAYGAN;	SANDOGLOBULIN;	GAMIMUNE)		,	•						-	•							,						

		No.
Preferred Indication Y	Cytomegalovirus disease	*
Exemplary Activity Assay		Neurite outgrowth assay, Neurosci Lett 2001 Mar 30;301(2):83-6 Cell adhesion assay, CAFCA, Centriftgal Assay for Fluorescence- based Cell Adhesion, Cancer Res 2001 Jan 1;61(1):339-47 Cell migration assay, Biochem Biophys Res Commun 2000 Nov 30;278(3):614-20
Biological Activity		Basement membrane protein; cell adhesion, differentiation, migration, signaling, neurite outgrowth and metastasis
PCT/Patent Reference		WO9506660 US5658789-A WO8911217-A2 WO9811217-A2 WO95111462-A WO9508628-A2 WO200066732-A2 WO9919348-A1 WO9919348-A1 WO90066730-A2 US7267564-A WO90066731-A2 US7267564-A WO200066731-A2 US5658789-A WO200058473-A2
Exemplary Identifier		Locus D: 3907 Locus D: 3908 Locus D: 3908 Locus D: 3909 Locus D: 3910 Locus D: 3911 Locus D: 3912 Locus D: 3913 Locus D: 3913 Locus D: 3913 Locus D: 3914 Locus D: 3913 Locus D: 3914 Locus D: 3913 Locus D: 3914 Locus D: 3913 Locus D: 3913 Locus D: 3913 INP 0002 18 INP 0022 81 INP 006 95 INP 00
Therapeutic Protein X	IVIG-CMV (Cytomegalovirus immune globulin intravenous (human); CMV IVIG;	m

Therapeutic Protein X		PCT/Patent	Biological Activity	Exemplary Activity Assay	Preferred Indication Y
	Identifier	Kelerence			
MCP/MCAF	LocusID:6347	US5714578-A ·	14578-A . Chemotactic factor for	Transendothelial lymphocyte	Cancer; Chemoprotection;
(Monocyte Chemotactic LocusID:6355	LocusID:6355	US7330446-A	monocytes; chemokine involved	monocytes; chemokine involved chemotaxis assay: Carr, M.W., et	Wounds
Protein: Monocyte	LocusID:6354	US6090795-A	in recruiting leukocytes during	al., Proc. Natl. Acad. Sci. USA,	
chemoattracting	_	US7304234-A	inflammation; attracts	vol. 91, pp. 3652-3656 (April	
neptides)		US5571713-A	macrophages during	1994).	
7	NP_006264	US5605671-A	inflammation and metastasis	-	
		WO9725427-A1	3		
	,	EP906954-A1			
- 3		WO9912968-A2	٠.		
		WO9509232-A			
-		EP488900-A			
-		WO9504158-A			
-	-	WO9509232-A			

	PCT/Patent Rinhogical Activity	Exemplary Activity Assay	Preferred Indication Y
CAS-148637-05-2 US5171675-A LocusID: 1435 NP 000748 US5573930-A WP 000748 US5681719-A US5681719-A US5643563-A US561150-A US561150-A US6117422-A US6103224-A US616851-A US6146851-A US6146851-A US6146851-A			
LocusID: 1435 US4929700-A NP 000748 US5573930-A XP 002150 US5681719-A US5843563-A US5861150-A US6117422-A US6103224-A US6146851-A US6146851-A US6146851-A US6146851-A	A-5791		Cancer; Hypercholesterolemia
XP_002150 US5672343-A US5681719-A US5861150-A US6117422-A US6103224-A US6146851-A US6146851-A Genbank::CAC20427 WO9931233-A1	US4929700-A macrophage/granulocyte-	tormation assay by the development of colonies containing macrophages (
US5681719-A US5681719-A US5643563-A US6103224-A US6103224-A US6146851-A US6146851-A	7343-A	-	
USS643563-A USS81150-A USS117422-A US6103224-A US6156300-A US6146851-A US6146851-A		proliferation and differentiation of Nov;2(6):356-67). M-CSF is also	
USS61150-A US6117422-A US6103224-A US6146851-A US6146851-A US6146851-A		detected in specific Bioassays with	
US6117422-A US6103224-A US6146851-A US6146851-A US6146851-A		cells lines that depend in their	
US6103224-A US6156300-A US6146851-A US6146851-A Genbank:CAC20427 WO9931233-A1			
US6156300-A US6146851-A US6146851-A Genbank:CAC20427 WO9931233-A1	-		*
US6146851-A Genbank:CAC20427 WO9931233-A1		ony example, BAC1.2F5; BaF3;	
Genbank:CAC20427 W09931233-A1			
Genbank:CAC20427 W09931233-A1	observes the phenomenon of	entirely different detection method is	
Genbank: CAC20427 W09931233-A1	synergistic suppression, i. e., the	the RT-PCR quantitation of cytokines.	
Genbank::CAC20427 W09931233-A1	combination of these two factors	ors	
Genbank: CAC20427 W09931233-A1	leads to a partial suppression of	Jc	-
Genbank::CAC20427 W09931233-A1	the generation of macrophage-		
Genbank::CAC20427 W09931233-A1	containing cell colonies. M-CSF	SSF	-
Genbank: CAC20427 W09931233-A1	is a specific factor in that the		
Genbank:CAC20427 W09931233-A1	proliferation inducing activity is	is	
Genbank: CAC20427 W09931233-A1	more or less restricted to the		•
Genbank: CAC20427 W09931233-A1	macrophage lineage. M-CSF also	osli	
Genbank:CAC20427 W09931233-A1	is a potent stimulator of	÷	
Genbank:CAC20427 W09931233-A1	functional activities of		
Genbank:CAC20427 W09931233-A1	monocytes. In normal human		
Genbank:CAC20427 W09931233-A1	macrophages M-CSF induces		
Genbank:CAC20427 W09931233-A1	antibody-dependent cellular	,	
Genbank:CAC20427 W09931233-A1	cytotoxicity. In monocytes and	Į pi	
Genbank:CAC20427 WO9931233-A1	macrophages M-CSF induces the	the	
Genbank:CAC20427 WO9931233-A1	synthesis of IL1, G-CSF, IFN,		
Genbank: CAC20427 WO9931233-A1	TNF, plasminogen activator,		
Genbank: CAC20427 WO9931233-A1	thromboplastin, prostaglandins and thromboxanes.		
ulating factor)		Transendothelial lymphocyte	Wound healing
		chemotaxis assay: Carr, M. W., et	
		vol. 91, pp. 3652-3656 (April	
		1994).	

Preferred Indication Y		Bone Fractures	Bleeding Disorders; Colorectal Cancer; Diabetic Retinopathy; Glioma; Heparin Neutralization after Cardiac Cartheterization or Cardiopulmonary Bypass Surgery; Kaposi's Sarcoma; Malignant Melanoma; Renal Cancer
Exemplary Activity Assay		Cell Attachment Assay: Senger DR, Perruzzi CA, Papadopoulos-Sergiou A, Van de Water L. (1994) Adhesive properties of osteopontin: regulation by a naturally occurring thrombin-cleavage in close proximity to the GRGDS cellbinding domain. Mol Biol Cell 5(5):565-74	Anti-angeogenic assay (PMID: 11259363)
Biological Activity	•	Osteopontin (OPN) is a highly phosphorylated sialoprotein that is a prominent component of the mineralized extracellular matrices of bones and teeth. OPN is characterized by the presence of a polyaspartic acid sequence and sites of Ser/Thr phosphorylation that mediate hydroxyapatite binding, and a highly conserved RGD motif that mediates cell attachment/signaling. Expression of OPN in a variety of tissues indicates a multiplicity of functions that involve one or more of more of more of a range of biological activities including developmental processes, wound healing, immunological responses, tumorigenesis, bone responses, tumorigenesis, bone	Platelet factor 4 (PF-4) is a CXC- Anti-angeogenic assay (PMID: chemokine with strong anti-angiogenic properties.
PCT/Patent Reference		WO9915904-A1 WO200062065- A1 WO9222316-A	W09302192-A W09504158-A US5248666-A US5776892-A
Exemplary Identifier		LocusID:6696 NP_000573 XP_011125	CAS-37270-94-3 LocusID:5196 NP_002610 XP_003505
Therapeutic Protein X	NCAF (Neutrophil chemoattracting	spl, ETA- Irotein 1, bone in 1, early T- re activation	Platelet Factor 4 (Endostatin B; Iroplact; RG 1001; Replistatin)

Therapeutic Protein X	Exemplary Identifier	PCT/Patent Reference	Biological Activity	Exemplary Activity Assay	Preferred Indication Y
Protein C (Drotrecogin CAS-60202 alfa; Activated Protein LocusID:56 C; CTC 111; Ceprotin; NP_000303 rhAPC; Zovant)	CAS-60202-16-6 LocusID:5624 NP_000303 XP_002706	WO9109953-A WO9112320-A US5516650-A US5358932-A	Protein C is a serine protease involved in coagulation and fibrinolysis.	Protein C activity may be assayed in vitro using a coagulation assay. (J Biol Chem 2000 Sep 1; 275(35): 27123-27128; Thromb Haemost 1994 Mar;71(3):339-346).	Disseminated intravascular coagulation; Septic shock; Thrombosis
Prothrombin, F2) Thrombin, F2)	LocusID:2147 NP_000497	WO9313208-A US5502034-A US6110721-A	Coagulation factor II is proteolytically cleaved to form thrombin in the first step of the coagulation cascade, which ultimately results in the stemming of blood loss. F2 also plays a role in maintaining vascular integrity.	Prothrombin quantitation and activation assay. "CA-1 method, a novel assay for quantification of normal prothrombin using a Ca2+dependent prothrombin activator, carinactivase-1." Thromb Res. 1999 May 15;94(4):221-6. "Activation of human prothrombin by argininespecific cysteine proteinases (Gingipains R) from Porphyromonas gingivalis." J Biol Chem. 2001 Mar 16	
Rabies IG (Rabies Immune Globulin; BAYRAB; HYPERRAB; IMOGAM RABIES-HT: IMOGAM)					Rabies
RhoD IG (RhoD Immune Globulin; IVIG-Rho(D); PAYRHO-D; MICRHOGAM; RHOGAM; WinRho SDF)		. ,			Prevention of isoimmunization of RhoD negative women at time of spontaneous or induced abortion or transfusion in pregnancy; Hemolyic disease; immune thrombocytopenic purpura; HIV infection
RSV IVIG (Respiratory syncytial virus IV immune globulin (human); Hypermune RSV; RESPIGAM)					Lower respiratory tract infections; respiratory syncytial infections

Preferred Indication Y		
Exemplary Activity Assay	BchE activity assay "Differential inhibition of human serum cholinesterase with fluoride: recognition of two new phenotypes." Nature 191: 496-498, 1961. "A rare genetically determined variant of pseudocholinesterase in two German families with high plasma enzyme activity." Europ. J. Biochem. 99: 65-69, 1979. "Genetic analysis of a Japanese patient with butyrylcholinesterase deficiency." Ann. Hum. Genet. 61:491-496, 1997.	cell adhesion assay. "Cell adhesion to fibronectin and tenascin: quantitative measurements of initial binding and subsequent strengthening response." J Cell Biol. 1989 Oct;109(4 Pt 1):1795-805.; "Tenascin interferes with fibronectin action." Cell. 1988 May 6;53(3):383-90. neurite growth in vitro. "Tenascin is accumulated along developing peripheral nerves and allows neurite outgrowth in vitro." Development. 1990 Oct;110(2):401-15.
Biological Activity	Also known as butyrylcholinesterase/pseuocholin esterase E1(CHE1). Human cholinesterase with fluoride tissues have two distinct cholinesterase activities: acetylcholinesterase and butyrylcholinesterase and transmission of nerve impulses, whereas the transmission of nerve impulses, whereas the transmission of butyrylcholinesterase remains unknown. Ann. Hum. Genet. 61:491-1997. BochE activity assay "Differ cognition of two new pherecognition of two new pherecognition of the muscle activity." Europ. J. Bioche impulses, whereas the collinesterase or the absence of its activity leads to prolonged apnea following administration of the muscle relaxant suxamethonium. The widespread expression of CHE1 in early differentiation suggests development-related functions for this protein.	The tenascins (TN) are a family of extracellular matrix proteins. The genes are expressed in distinct tissues at different times during embyronic development and are present in adult tissues. TN-R is detected predominantly in the central nervous system of early embryos and likely involved in central nervous system development. TN-XA is overexpressed in many tumors.
PCT/Patent Reference	WO9107483-A WO9523158-A US6001625-A US5695750-A	WO9628550-A1 WO9608513-A1 US5681931-A US5635360-A WO9222319-A WO9608513-A1 US5681931-A US5635360-A US5635360-A
. Exemplary Identifier	LocusID:590 LocusID:1110 NP_000046 XP_003134	LocusID:7143 LocusID:7146 LocusID:7148 NP_003276 NP_009047 XP_001730 XP_004201
Therapeutic Protein X	Serum Cholinesterase	Tenascin

Preferred Indication V	Tetanus ·	Atherosclerosis; Vascular Restenosis; Cancer; Cardiovascular Disorders; Malignant Melanoma; Clotting disorders; Transplantation.
Exemplary Activity Assay		Individual functions of the molecule are assayed separately. The cell adhesion function is assayed using scell adhesion assay (Feinberg and Vogelstein, 1983; Anal. Biochem. 132 pp6-10); PAI binding using a solid phase binding assay (Seiffert & Loskutoff, 1991; J. Biol. Chem. 266 pp2824-2830)
Biological Activity		Binds to serpin serine protease inhibitors such as PAI, mediates cell-to-substrate adhesion, inhibits the cytolytic action of the terminal complement cascade in vitro and inhibits inactivation of thrombin by antithrombin, thereby regulating coagulation.
PCT/Patent Reference		US514582-A US6140072-A WO9213075-A
Exemplary Identifier		LocusID:7448 NP_000629 XP_008484
Therapeutic Protein X	Tetanus IG (Tetanus Immune Globulin; TIG; BAYTET)	

In preferred embodiments, the albumin fusion proteins of the invention are capable of a therapeutic activity and/or biologic activity corresponding to the therapeutic activity and/or biologic activity of the Therapeutic protein corresponding to the Therapeutic protein portion of the albumin fusion protein listed in the corresponding row of Table 1. (See, e.g., the "Biological Activity" and "Therapeutic Protein X"columns of Table 1.) In further preferred embodiments, the therapeutically active protein portions of the albumin fusion proteins of the invention are fragments or variants of the reference sequence cited in the "Exemplary Identifier" column of Table 1, and are capable of the therapeutic activity and/or biologic activity of the corresponding Therapeutic protein disclosed in "Biological Activity" column of Table 1.

Polypeptide and Polynucleotide Fragments and Variants

Fragments

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The present invention is further directed to fragments of the Therapeutic proteins described in Table 1, albumin proteins, and/or albumin fusion proteins of the invention.

Even if deletion of one or more amino acids from the N-terminus of a protein results in modification or loss of one or more biological functions of the Therapeutic protein, albumin protein, and/or albumin fusion protein, other Therapeutic activities and/or functional activities (e.g., biological activities, ability to multimerize, ability to bind a ligand) may still be retained. For example, the ability of polypeptides with N-terminal deletions to induce and/or bind to antibodies which recognize the complete or mature forms of the polypeptides generally will be retained when less than the majority of the residues of the complete polypeptide are removed from the N-terminus. Whether a particular polypeptide lacking N-terminal residues of a complete polypeptide retains such immunologic activities can readily be determined by routine methods described herein and otherwise known in the art. It is not unlikely that a mutein with a large number of deleted N-terminal amino acid residues may retain some biological or immunogenic activities. In fact, peptides composed of as few as six amino acid residues may often evoke an immune response.

Accordingly, fragments of a Therapeutic protein corresponding to a Therapeutic protein portion of an albumin fusion protein of the invention, include the full length protein as well as polypeptides having one or more residues deleted from the amino terminus of the amino acid sequence of the reference polypeptide (e.g., a Therapeutic protein as disclosed in Table 1). In particular, N-terminal deletions may be described by the general formula m-q, where q is a whole integer representing the total number of amino acid residues in a reference polypeptide (e.g., a Therapeutic protein referred to in Table 1), and m is defined as any integer ranging from 2 to q-6. Polynucleotides encoding these polypeptides are also

encompassed by the invention.

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In addition, fragments of serum albumin polypeptides corresponding to an albumin protein portion of an albumin fusion protein of the invention, include the full length protein as well as polypeptides having one or more residues deleted from the amino terminus of the amino acid sequence of the reference polypeptide (i.e., serum albumin). In particular, N-terminal deletions may be described by the general formula m-585, where 585 is a whole integer representing the total number of amino acid residues in serum albumin (SEQ ID NO:18), and m is defined as any integer ranging from 2 to 579. Polynucleotides encoding these polypeptides are also encompassed by the invention.

Moreover, fragments of albumin fusion proteins of the invention, include the full length albumin fusion protein as well as polypeptides having one or more residues deleted from the amino terminus of the albumin fusion protein. In particular, N-terminal deletions may be described by the general formula m-q, where q is a whole integer representing the total number of amino acid residues in the albumin fusion protein, and m is defined as any integer ranging from 2 to q-6. Polynucleotides encoding these polypeptides are also encompassed by the invention.

Also as mentioned above, even if deletion of one or more amino acids from the N-terminus or C-terminus of a reference polypeptide (e.g., a Therapeutic protein and/or serum albumin protein) results in modification or loss of one or more biological functions of the protein, other functional activities (e.g., biological activities, ability to multimerize, ability to bind a ligand) and/or Therapeutic activities may still be retained. For example the ability of polypeptides with C-terminal deletions to induce and/or bind to antibodies which recognize the complete or mature forms of the polypeptide generally will be retained when less than the majority of the residues of the complete or mature polypeptide are removed from the C-terminus. Whether a particular polypeptide lacking the N-terminal and/or C-terminal residues of a reference polypeptide retains Therapeutic activity can readily be determined by routine methods described herein and/or otherwise known in the art.

The present invention further provides polypeptides having one or more residues deleted from the carboxy terminus of the amino acid sequence of a Therapeutic protein corresponding to a Therapeutic protein portion of an albumin fusion protein of the invention (e.g., a Therapeutic protein referred to in Table 1). In particular, C-terminal deletions may be described by the general formula 1-n, where n is any whole integer ranging from 6 to q-1, and where q is a whole integer representing the total number of amino acid residues in a reference polypeptide (e.g., a Therapeutic protein referred to in Table 1). Polynucleotides encoding these polypeptides are also encompassed by the invention.

In addition, the present invention provides polypeptides having one or more residues deleted from the carboxy terminus of the amino acid sequence of an albumin protein

corresponding to an albumin protein portion of an albumin fusion protein of the invention (e.g., serum albumin). In particular, C-terminal deletions may be described by the general formula 1-n, where n is any whole integer ranging from 6 to 584, where 584 is the whole integer representing the total number of amino acid residues in serum albumin (SEQ ID NO:18) minus 1. Polynucleotides encoding these polypeptides are also encompassed by the invention.

Moreover, the present invention provides polypeptides having one or more residues deleted from the carboxy terminus of an albumin fusion protein of the invention. In particular, C-terminal deletions may be described by the general formula 1-n, where n is any whole integer ranging from 6 to q-1, and where q is a whole integer representing the total number of amino acid residues in an albumin fusion protein of the invention. Polynucleotides encoding these polypeptides are also encompassed by the invention.

In addition, any of the above described N- or C-terminal deletions can be combined to produce a N- and C-terminal deleted reference polypeptide. The invention also provides polypeptides having one or more amino acids deleted from both the amino and the carboxyl termini, which may be described generally as having residues m-n of a reference polypeptide (e.g., a Therapeutic protein referred to in Table 1, or serum albumin (e.g., SEQ ID NO:18), or an albumin fusion protein of the invention) where n and m are integers as described above. Polynucleotides encoding these polypeptides are also encompassed by the invention.

The present application is also directed to proteins containing polypeptides at least 80%, 85%, 90%, 95%, 96%, 97%, 98% or 99% identical to a reference polypeptide sequence (e.g., a Therapeutic protein, serum albumin protein or an albumin fusion protein of the invention) set forth herein, or fragments thereof. In preferred embodiments, the application is directed to proteins comprising polypeptides at least 80%, 85%, 90%, 95%, 96%, 97%, 98% or 99% identical to reference polypeptides having the amino acid sequence of N- and C-terminal deletions as described above. Polynucleotides encoding these polypeptides are also encompassed by the invention.

Preferred polypeptide fragments of the invention are fragments comprising, or alternatively, consisting of, an amino acid sequence that displays a Therapeutic activity and/or functional activity (e.g. biological activity) of the polypeptide sequence of the Therapeutic protein or serum albumin protein of which the amino acid sequence is a fragment.

Other preferred polypeptide fragments are higherically active fragments. Piclogically active

Other preferred polypeptide fragments are biologically active fragments. Biologically active fragments are those exhibiting activity similar, but not necessarily identical, to an activity of the polypeptide of the present invention. The biological activity of the fragments may include an improved desired activity, or a decreased undesirable activity.

Variants

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"Variant" refers to a polynucleotide or nucleic acid differing from a reference nucleic acid or polypeptide, but retaining essential properties thereof. Generally, variants are overall closely similar, and, in many regions, identical to the reference nucleic acid or polypeptide.

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As used herein, "variant", refers to a Therapeutic protein portion of an albumin fusion protein of the invention, albumin portion of an albumin fusion protein of the invention, or albumin fusion protein differing in sequence from a Therapeutic protein (e.g. see "therapeutic" column of Table 1), albumin protein, and/or albumin fusion protein of the invention, respectively, but retaining at least one functional and/or therapeutic property thereof (e.g., a therapeutic activity and/or biological activity as disclosed in the "Biological Activity" column of Table 1) as described elsewhere herein or otherwise known in the art. Generally, variants are overall very similar, and, in many regions, identical to the amino acid sequence of the Therapeutic protein corresponding to a Therapeutic protein portion of an albumin fusion protein of the invention, albumin protein corresponding to an albumin protein portion of an albumin fusion protein of the invention, and/or albumin fusion protein of the invention. Nucleic acids encoding these variants are also encompassed by the invention.

The present invention is also directed to proteins which comprise, or alternatively consist of, an amino acid sequence which is at least 80%, 85%, 90%, 95%, 96%, 97%, 98%, 99% or 100%, identical to, for example, the amino acid sequence of a Therapeutic protein corresponding to a Therapeutic protein portion of an albumin fusion protein of the invention (e.g., an amino acid sequence disclosed in the "Exemplary Identifier" column of Table 1, or fragments or variants thereof), albumin proteins (e.g., SEQ ID NO:18 or fragments or variants thereof) corresponding to an albumin protein portion of an albumin fusion protein of the invention, and/or albumin fusion proteins of the invention. Fragments of these polypeptides are also provided (e.g., those fragments described herein). Further polypeptides encompassed by the invention are polypeptides encoded by polynucleotides which hybridize to the complement of a nucleic acid molecule encoding an amino acid sequence of the invention under stringent hybridization conditions (e.g., hybridization to filter bound DNA in 6X Sodium chloride/Sodium citrate (SSC) at about 45 degrees Celsius. followed by one or more washes in 0.2X SSC, 0.1% SDS at about 50 - 65 degrees Celsius), under highly stringent conditions (e.g., hybridization to filter bound DNA in 6X sodium chloride/Sodium citrate (SSC) at about 45 degrees Celsius, followed by one or more washes in 0.1X SSC, 0.2% SDS at about 68 degrees Celsius), or under other stringent hybridization conditions which are known to those of skill in the art (see, for example, Ausubel, F.M. et al., eds., 1989 Current protocol in Molecular Biology, Green publishing associates, Inc., and John Wiley & Sons Inc., New York, at pages 6.3.1 - 6.3.6 and 2.10.3). Polynucleotides encoding these polypeptides are also encompassed by the invention.

By a polypeptide having an amino acid sequence at least, for example, 95% "identical"

to a query amino acid sequence of the present invention, it is intended that the amino acid sequence of the subject polypeptide is identical to the query sequence except that the subject polypeptide sequence may include up to five amino acid alterations per each 100 amino acids of the query amino acid sequence. In other words, to obtain a polypeptide having an amino acid sequence at least 95% identical to a query amino acid sequence, up to 5% of the amino acid residues in the subject sequence may be inserted, deleted, or substituted with another amino acid. These alterations of the reference sequence may occur at the amino- or carboxy-terminal positions of the reference amino acid sequence or anywhere between those terminal positions, interspersed either individually among residues in the reference sequence or in one or more contiguous groups within the reference sequence.

As a practical matter, whether any particular polypeptide is at least 80%, 85%, 90%, 95%, 96%, 97%, 98% or 99% identical to, for instance, the amino acid sequence of an albumin fusion protein of the invention or a fragment thereof (such as the Therapeutic protein portion of the albumin fusion protein or the albumin portion of the albumin fusion protein), can be determined conventionally using known computer programs. A preferred method for determining the best overall match between a query sequence (a sequence of the present invention) and a subject sequence, also referred to as a global sequence alignment, can be determined using the FASTDB computer program based on the algorithm of Brutlag et al. (Comp. App. Biosci.6:237-245 (1990)). In a sequence alignment the query and subject sequences are either both nucleotide sequences or both amino acid sequences. The result of said global sequence alignment is expressed as percent identity. Preferred parameters used in a FASTDB amino acid alignment are: Matrix=PAM 0, k-tuple=2, Mismatch Penalty=1, Joining Penalty=20, Randomization Group Length=0, Cutoff Score=1, Window Size=sequence length, Gap Penalty=5, Gap Size Penalty=0.05, Window Size=500 or the length of the subject amino acid sequence, whichever is shorter.

If the subject sequence is shorter than the query sequence due to N- or C-terminal deletions, not because of internal deletions, a manual correction must be made to the results. This is because the FASTDB program does not account for N- and C-terminal truncations of the subject sequence when calculating global percent identity. For subject sequences truncated at the N- and C-termini, relative to the query sequence, the percent identity is corrected by calculating the number of residues of the query sequence that are N- and C-terminal of the subject sequence, which are not matched/aligned with a corresponding subject residue, as a percent of the total bases of the query sequence. Whether a residue is matched/aligned is determined by results of the FASTDB sequence alignment. This percentage is then subtracted from the percent identity, calculated by the above FASTDB program using the specified parameters, to arrive at a final percent identity score. This final percent identity score is what is used for the purposes of the present invention. Only residues

to the N- and C-termini of the subject sequence, which are not matched/aligned with the query sequence, are considered for the purposes of manually adjusting the percent identity score. That is, only query residue positions outside the farthest N- and C- terminal residues of the subject sequence.

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For example, a 90 amino acid residue subject sequence is aligned with a 100 residue query sequence to determine percent identity. The deletion occurs at the N-terminus of the subject sequence and therefore, the FASTDB alignment does not show a matching/alignment of the first 10 residues at the N-terminus. The 10 unpaired residues represent 10% of the sequence (number of residues at the N- and C- termini not matched/total number of residues in the query sequence) so 10% is subtracted from the percent identity score calculated by the FASTDB program. If the remaining 90 residues were perfectly matched the final percent identity would be 90%. In another example, a 90 residue subject sequence is compared with a 100 residue query sequence. This time the deletions are internal deletions so there are no residues at the N- or C-termini of the subject sequence which are not matched/aligned with the query. In this case the percent identity calculated by FASTDB is not manually corrected. Once again, only residue positions outside the N- and C-terminal ends of the subject sequence, as displayed in the FASTDB alignment, which are not matched/aligned with the query sequence are manually corrected for. No other manual corrections are to made for the purposes of the present invention.

The variant will usually have at least 75 % (preferably at least about 80%, 90%, 95% or 99%) sequence identity with a length of normal HA or Therapeutic protein which is the same length as the variant. Homology or identity at the nucleotide or amino acid sequence level is determined by BLAST (Basic Local Alignment Search Tool) analysis using the algorithm employed by the programs blastp, blastn, blastx, tblastn and tblastx (Karlin *et al.*, Proc. Natl. Acad. Sci. USA 87: 2264-2268 (1990) and Altschul, J. Mol. Evol. 36: 290-300 (1993), fully incorporated by reference) which are tailored for sequence similarity searching.

The approach used by the BLAST program is to first consider similar segments between a query sequence and a database sequence, then to evaluate the statistical significance of all matches that are identified and finally to summarize only those matches which satisfy a preselected threshold of significance. For a discussion of basic issues in similarity searching of sequence databases, see Altschul et al., (Nature Genetics 6: 119-129 (1994)) which is fully incorporated by reference. The search parameters for histogram, descriptions, alignments, expect (i.e., the statistical significance threshold for reporting matches against database sequences), cutoff, matrix and filter are at the default settings. The default scoring matrix used by blastp, blastx, tblastn, and tblastx is the BLOSUM62 matrix (Henikoff et al., Proc. Natl. Acad. Sci. USA 89: 10915-10919 (1992), fully incorporated by reference). For blastn, the scoring matrix is set by the ratios of M (i.e., the reward score for a pair of matching residues)

to N (i.e., the penalty score for mismatching residues), wherein the default values for M and N are 5 and -4, respectively. Four blastn parameters may be adjusted as follows: Q=10 (gap creation penalty); R=10 (gap extension penalty); wink=1 (generates word hits at every winkth position along the query); and gapw=16 (sets the window width within which gapped alignments are generated). The equivalent Blastp parameter settings were Q=9; R=2; wink=1; and gapw=32. A Bestfit comparison between sequences, available in the GCG package version 10.0, uses DNA parameters GAP=50 (gap creation penalty) and LEN=3 (gap extension penalty) and the equivalent settings in protein comparisons are GAP=8 and LEN=2.

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The polynucleotide variants of the invention may contain alterations in the coding regions, non-coding regions, or both. Especially preferred are polynucleotide variants containing alterations which produce silent substitutions, additions, or deletions, but do not alter the properties or activities of the encoded polypeptide. Nucleotide variants produced by silent substitutions due to the degeneracy of the genetic code are preferred. Moreover, polypeptide variants in which less than 50, less than 40, less than 30, less than 20, less than 10, or 5-50, 5-25, 5-10, 1-5, or 1-2 amino acids are substituted, deleted, or added in any combination are also preferred. Polynucleotide variants can be produced for a variety of reasons, e.g., to optimize codon expression for a particular host (change codons in the human mRNA to those preferred by a bacterial host, such as, yeast or *E. coli*).

In a preferred embodiment, a polynucleotide encoding an albumin portion of an albumin fusion protein of the invention is optimized for expression in yeast or mammalian cells. In further preferred embodiment, a polynucleotide encoding a Therapeutic protein portion of an albumin fusion protein of the invention is optimized for expression in yeast or mammalian cells. In a still further preferred embodiment, a polynucleotide encoding an albumin fusion protein of the invention is optimized for expression in yeast or mammalian cells.

In an alternative embodiment, a codon optimized polynucleotide encoding a Therapeutic protein portion of an albumin fusion protein of the invention does not hybridize to the wild type polynucleotide encoding the Therapeutic protein under stringent hybridization conditions as described herein. In a further embodiment, a codon optimized polynucleotide encoding an albumin portion of an albumin fusion protein of the invention does not hybridize to the wild type polynucleotide encoding the albumin protein under stringent hybridization conditions as described herein. In another embodiment, a codon optimized polynucleotide encoding an albumin fusion protein of the invention does not hybridize to the wild type polynucleotide encoding the Therapeutic protein portin or the albumin protein portion under stringent hybridization conditions as described herein.

In an additional embodiment, polynucleotides encoding a Therapeutic protein portion of an albumin fusion protein of the invention do not comprise, or alternatively consist of, the

naturally occurring sequence of that Therapeutic protein. In a further embodiment, polynucleotides encoding an albumin protein portion of an albumin fusion protein of the invention do not comprise, or alternatively consist of, the naturally occurring sequence of albumin protein. In an alternative embodiment, polynucleotides encoding an albumin fusion protein of the invention do not comprise, or alternatively consist of, the naturally occurring sequence of a Therapeutic protein portion or the albumin protein portion.

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Naturally occurring variants are called "allelic variants," and refer to one of several alternate forms of a gene occupying a given locus on a chromosome of an organism. (Genes II, Lewin, B., ed., John Wiley & Sons, New York (1985)). These allelic variants can vary at either the polynucleotide and/or polypeptide level and are included in the present invention. Alternatively, non-naturally occurring variants may be produced by mutagenesis techniques or by direct synthesis.

Using known methods of protein engineering and recombinant DNA technology, variants may be generated to improve or alter the characteristics of the polypeptides of the present invention. For instance, one or more amino acids can be deleted from the N-terminus or C-terminus of the polypeptide of the present invention without substantial loss of biological function. As an example, Ron et al. (J. Biol. Chem. 268: 2984-2988 (1993)) reported variant KGF proteins having heparin binding activity even after deleting 3, 8, or 27 amino-terminal amino acid residues. Similarly, Interferon gamma exhibited up to ten times higher activity after deleting 8-10 amino acid residues from the carboxy terminus of this protein. (Dobeli et al., J. Biotechnology 7:199-216 (1988).)

Moreover, ample evidence demonstrates that variants often retain a biological activity similar to that of the naturally occurring protein. For example, Gayle and coworkers (J. Biol. Chem. 268:22105-22111 (1993)) conducted extensive mutational analysis of human cytokine IL-1a. They used random mutagenesis to generate over 3,500 individual IL-1a mutants that averaged 2.5 amino acid changes per variant over the entire length of the molecule. Multiple mutations were examined at every possible amino acid position. The investigators found that "[m]ost of the molecule could be altered with little effect on either [binding or biological activity]." In fact, only 23 unique amino acid sequences, out of more than 3,500 nucleotide sequences examined, produced a protein that significantly differed in activity from wild-type.

Furthermore, even if deleting one or more amino acids from the N-terminus or C-terminus of a polypeptide results in modification or loss of one or more biological functions, other biological activities may still be retained. For example, the ability of a deletion variant to induce and/or to bind antibodies which recognize the secreted form will likely be retained when less than the majority of the residues of the secreted form are removed from the N-terminus or C-terminus. Whether a particular polypeptide lacking N- or C-terminal residues of a protein retains such immunogenic activities can readily be determined by routine methods

described herein and otherwise known in the art.

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Thus, the invention further includes polypeptide variants which have a functional activity (e.g., biological activity and/or therapeutic activity). In highly preferred embodiments the invention provides variants of albumin fusion proteins that have a functional activity (e.g., biological activity and/or therapeutic activity, such as that disclosed in the "Biological Activity" column in Table 1) that corresponds to one or more biological and/or therapeutic activities of the Therapeutic protein corresponding to the Therapeutic protein portion of the albumin fusion protein. Such variants include deletions, insertions, inversions, repeats, and substitutions selected according to general rules known in the art so as have little effect on activity.

In preferred embodiments, the variants of the invention have conservative substitutions. By "conservative substitutions" is intended swaps within groups such as replacement of the aliphatic or hydrophobic amino acids Ala, Val, Leu and Ile; replacement of the hydroxyl residues Ser and Thr; replacement of the acidic residues Asp and Glu; replacement of the amide residues Asn and Gln, replacement of the basic residues Lys, Arg, and His; replacement of the aromatic residues Phe, Tyr, and Trp, and replacement of the small-sized amino acids Ala, Ser, Thr, Met, and Gly.

Guidance concerning how to make phenotypically silent amino acid substitutions is provided, for example, in Bowie et al., "Deciphering the Message in Protein Sequences: Tolerance to Amino Acid Substitutions," Science 247:1306-1310 (1990), wherein the authors indicate that there are two main strategies for studying the tolerance of an amino acid sequence to change.

The first strategy exploits the tolerance of amino acid substitutions by natural selection during the process of evolution. By comparing amino acid sequences in different species, conserved amino acids can be identified. These conserved amino acids are likely important for protein function. In contrast, the amino acid positions where substitutions have been tolerated by natural selection indicates that these positions are not critical for protein function. Thus, positions tolerating amino acid substitution could be modified while still maintaining biological activity of the protein.

The second strategy uses genetic engineering to introduce amino acid changes at specific positions of a cloned gene to identify regions critical for protein function. For example, site directed mutagenesis or alanine-scanning mutagenesis (introduction of single alanine mutations at every residue in the molecule) can be used. See Cunningham and Wells, Science 244:1081-1085 (1989). The resulting mutant molecules can then be tested for biological activity.

As the authors state, these two strategies have revealed that proteins are surprisingly tolerant of amino acid substitutions. The authors further indicate which amino acid changes

are likely to be permissive at certain amino acid positions in the protein. For example, most buried (within the tertiary structure of the protein) amino acid residues require nonpolar side chains, whereas few features of surface side chains are generally conserved. tolerated conservative amino acid substitutions involve replacement of the aliphatic or hydrophobic amino acids Ala, Val, Leu and Ile; replacement of the hydroxyl residues Ser and Thr; replacement of the acidic residues Asp and Glu; replacement of the amide residues Asn and Gln. replacement of the basic residues Lys, Arg, and His; replacement of the aromatic residues Phe, Tyr, and Trp, and replacement of the small-sized amino acids Ala, Ser, Thr. Met, and Gly. Besides conservative amino acid substitution, variants of the present invention include (i) polypeptides containing substitutions of one or more of the non-conserved amino acid residues, where the substituted amino acid residues may or may not be one encoded by the genetic code, or (ii) polypeptides containing substitutions of one or more of the amino acid residues having a substituent group, or (iii) polypeptides which have been fused with or chemically conjugated to another compound, such as a compound to increase the stability and/or solubility of the polypeptide (for example, polyethylene glycol), (iv) polypeptide containing additional amino acids, such as, for example, an IgG Fc fusion region peptide, . Such variant polypeptides are deemed to be within the scope of those skilled in the art from the teachings herein.

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For example, polypeptide variants containing amino acid substitutions of charged amino acids with other charged or neutral amino acids may produce proteins with improved characteristics, such as less aggregation. Aggregation of pharmaceutical formulations both reduces activity and increases clearance due to the aggregate's immunogenic activity. See Pinckard et al., Clin. Exp. Immunol. 2:331-340 (1967); Robbins et al., Diabetes 36: 838-845 (1987); Cleland et al., Crit. Rev. Therapeutic Drug Carrier Systems 10:307-377 (1993).

In specific embodiments, the polypeptides of the invention comprise; or alternatively, consist of, fragments or variants of the amino acid sequence of a Therapeutic protein described herein and/or human serum albumin, and/or albumin fusion protein of the invention, wherein the fragments or variants have 1-5, 5-10, 5-25, 5-50, 10-50 or 50-150, amino acid residue additions, substitutions, and/or deletions when compared to the reference amino acid sequence. In preferred embodiments, the amino acid substitutions are conservative. Nucleic acids encoding these polypeptides are also encompassed by the invention.

The polypeptide of the present invention can be composed of amino acids joined to each other by peptide bonds or modified peptide bonds, i.e., peptide isosteres, and may contain amino acids other than the 20 gene-encoded amino acids. The polypeptides may be modified by either natural processes, such as post-translational processing, or by chemical modification techniques which are well known in the art. Such modifications are well

described in basic texts and in more detailed monographs, as well as in a voluminous research Modifications can occur anywhere in a polypeptide, including the peptide backbone, the amino acid side-chains and the amino or carboxyl termini. It will be appreciated that the same type of modification may be present in the same or varying degrees at several sites in a given polypeptide. Also, a given polypeptide may contain many types of modifications. Polypeptides may be branched, for example, as a result of ubiquitination, and they may be cyclic, with or without-branching. Cyclic, branched, and branched cyclic polypeptides may result from posttranslation natural processes or may be made by synthetic methods. Modifications include acetylation, acylation, ADP-ribosylation, amidation, covalent attachment of flavin, covalent attachment of a heme moiety, covalent attachment of a nucleotide or nucleotide derivative, covalent attachment of a lipid or lipid derivative, covalent attachment of phosphotidylinositol, cross-linking, cyclization, disulfide bond formation, demethylation, formation of covalent cross-links, formation of cysteine, formation of pyroglutamate, formylation, gamma-carboxylation, glycosylation, GPI anchor formation, hydroxylation, iodination, methylation, myristylation, oxidation, pegylation, proteolytic processing, phosphorylation, prenylation, racemization, selenoylation, sulfation, transfer-RNA mediated addition of amino acids to proteins such as arginylation, and ubiquitination. (See, for instance, PROTEINS - STRUCTURE AND MOLECULAR PROPERTIES, 2nd Creighton, W. H. Freeman and Company, New York (1993); POST-TRANSLATIONAL COVALENT MODIFICATION OF PROTEINS, B. C. Johnson, Ed., Academic Press, New York, pgs. 1-12 (1983); Seifter et al., Meth. Enzymol. 182:626-646 (1990); Rattan et al., Ann. N.Y. Acad. Sci. 663:48-62 (1992)).

Functional activity

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"A polypeptide having functional activity" refers to a polypeptide capable of displaying one or more known functional activities associated with the full-length, proprotein, and/or mature form of a Therapeutic protein. Such functional activities include, but are not limited to, biological activity, antigenicity [ability to bind (or compete with a polypeptide for binding) to an anti-polypeptide antibody], immunogenicity (ability to generate antibody which binds to a specific polypeptide of the invention), ability to form multimers with polypeptides of the invention, and ability to bind to a receptor or ligand for a polypeptide.

"A polypeptide having biological activity" refers to a polypeptide exhibiting activity similar to, but not necessarily identical to, an activity of a Therapeutic protein of the present invention, including mature forms, as measured in a particular biological assay, with or without dose dependency. In the case where dose dependency does exist, it need not be identical to that of the polypeptide, but rather substantially similar to the dose-dependence in a

given activity as compared to the polypeptide of the present invention (i.e., the candidate polypeptide will exhibit greater activity or not more than about 25-fold less and, preferably, not more than about tenfold less activity, and most preferably, not more than about three-fold less activity relative to the polypeptide of the present invention).

In preferred embodiments, an albumin fusion protein of the invention has at least one biological and/or therapeutic activity associated with the Therapeutic protein (or fragment or variant thereof) when it is not fused to albumin.

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The albumin fusion proteins of the invention can be assayed for functional activity (e.g., biological activity) using or routinely modifying assays known in the art, as well as assays described herein. Specifically, albumin fusion proteins may be assayed for functional activity (e.g., biological activity or therapeutic activity) using the assay referenced in the "Exemplary Activity Assay" column of Table 1. Additionally, one of skill in the art may routinely assay fragments of a Therapeutic protein corresponding to a Therapeutic protein portion of an albumin fusion protein of the invention, for activity using assays referenced in its corresponding row of Table 1. Further, one of skill in the art may routinely assay fragments of an albumin protein corresponding to an albumin protein portion of an albumin fusion protein of the invention, for activity using assays known in the art and/or as described in the Examples section below.

For example, in one embodiment where one is assaying for the ability of an albumin fusion protein of the invention to bind or compete with a Therapeutic protein for binding to antibody and/or anti-albumin antibody, various anti-Therapeutic polypeptide immunoassays known in the art can be used, including but not limited to, competitive and non-competitive assay systems using techniques such as radioimmunoassays, ELISA (enzyme linked immunosorbent assay), "sandwich" immunoassays, immunoradiometric assays, gel diffusion precipitation reactions, immunodiffusion assays, in situ immunoassays (using colloidal gold, enzyme or radioisotope labels, for example), western blots, precipitation reactions, agglutination assays (e.g., gel agglutination assays, hemagglutination assays), complement fixation assays, immunofluorescence assays, protein A assays, and immunoelectrophoresis assays, etc. In one embodiment, antibody binding is detected by detecting a label on the primary antibody. In another embodiment, the primary antibody is detected by detecting binding of a secondary antibody or reagent to the primary antibody. In a further embodiment, the secondary antibody is labeled. Many means are known in the art for detecting binding in an immunoassay and are within the scope of the present invention.

In a preferred embodiment, where a binding partner (e.g., a receptor or a ligand) of a Therapeutic protein is identified, binding to that binding partner by an albumin fusion protein containing that Therapeutic protein as the Therapeutic protein portion of the fusion can be assayed, e.g., by means well-known in the art, such as, for example, reducing and non-

reducing gel chromatography, protein affinity chromatography, and affinity blotting. See generally, Phizicky et al., Microbiol. Rev. 59:94-123 (1995). In another embodiment, the ability of physiological correlates of an albumin fusion protein of the present invention to bind to a substrate(s) of the Therapeutic polypeptide corresponding to the Therapeutic portion of the albumin fusion protein of the invention can be routinely assayed using techniques known in the art.

In an alternative embodiment, where the ability of an albumin fusion protein of the invention to multimerize is being evaluated, association with other components of the multimer can be assayed, e.g., by means well-known in the art, such as, for example, reducing and non-reducing gel chromatography, protein affinity chromatography, and affinity blotting. See generally, Phizicky et al., *supra*.

In addition, assays described herein (see Examples and Table 1) and otherwise known in the art may routinely be applied to measure the ability of albumin fusion proteins of the present invention and fragments, variants and derivatives thereof to elicit biological activity and/or Therapeutic activity (either *in vitro* or *in vivo*) related to either the Therapeutic protein portion and/or albumin portion of the albumin fusion protein of the present invention. Other methods will be known to the skilled artisan and are within the scope of the invention.

Albumin

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As described above, an albumin fusion protein of the invention comprises at least a fragment or variant of a Therapeutic protein and at least a fragment or variant of human serum albumin, which are associated with one another, preferably by genetic fusion or chemical conjugation.

The terms, human serum albumin (HSA) and human albumin (HA) are used interchangeably herein. The terms, "albumin and "serum albumin" are broader, and encompass human serum albumin (and fragments and variants thereof) as well as albumin from other species (and fragments and variants thereof).

As used herein, "albumin" refers collectively to albumin protein or amino acid sequence, or an albumin fragment or variant, having one or more functional activities (e.g., biological activities) of albumin. In particular, "albumin" refers to human albumin or fragments thereof (see EP 201 239, EP 322 094 WO 97/24445, WO95/23857) especially the mature form of human albumin as shown in Figure 15 and SEQ ID NO:18, or albumin from other vertebrates or fragments thereof, or analogs or variants of these molecules or fragments thereof.

In preferred embodiments, the human serum albumin protein used in the albumin fusion proteins of the invention contains one or both of the following sets of point mutations with reference to SEQ ID NO:18: Leu-407 to Ala, Leu-408 to Val, Val-409 to Ala; and Arg-

410 to Ala; or Arg-410 to A, Lys-413 to Gln, and Lys-414 to Gln (see, e.g., International Publication No. WO95/23857, hereby incorporated in its entirety by reference herein). In even more preferred embodiments, albumin fusion proteins of the invention that contain one or both of above-described sets of point mutations have improved stability/resistance to yeast Yap3p proteolytic cleavage, allowing increased production of recombinant albumin fusion proteins expressed in yeast host cells.

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As used herein, a portion of albumin sufficient to prolong the therapeutic activity or shelf-life of the Therapeutic protein refers to a portion of albumin sufficient in length or structure to stabilize or prolong the therapeutic activity of the protein so that the shelf life of the Therapeutic protein portion of the albumin fusion protein is prolonged or extended compared to the shelf-life in the non-fusion state. The albumin portion of the albumin fusion proteins may comprise the full length of the HA sequence as described above or as shown in Figure 15, or may include one or more fragments thereof that are capable of stabilizing or prolonging the therapeutic activity. Such fragments may be of 10 or more amino acids in length or may include about 15, 20, 25, 30, 50, or more contiguous amino acids from the HA sequence or may include part or all of specific domains of HA. For instance, one or more fragments of HA spanning the first two immunoglobulin-like domains may be used.

The albumin portion of the albumin fusion proteins of the invention may be a variant of normal HA. The Therapeutic protein portion of the albumin fusion proteins of the invention may also be variants of the Therapeutic proteins as described herein. The term "variants" includes insertions, deletions and substitutions, either conservative or non conservative, where such changes do not substantially alter one or more of the oncotic, useful ligand-binding and non-immunogenic properties of albumin, or the active site, or active domain which confers the therapeutic activities of the Therapeutic proteins.

In particular, the albumin fusion proteins of the invention may include naturally occurring polymorphic variants of human albumin and fragments of human albumin, for example those fragments disclosed in EP 322 094 (namely HA (Pn), where n is 369 to 419). The albumin may be derived from any vertebrate, especially any mammal, for example human, cow, sheep, or pig. Non-mammalian albumins include, but are not limited to, hen and salmon. The albumin portion of the albumin fusion protein may be from a different animal than the Therapeutic protein portion.

Generally speaking, an HA fragment or variant will be at least 100 amino acids long, preferably at least 150 amino acids long. The HA variant may consist of or alternatively comprise at least one whole domain of HA, for example domains 1 (amino acids 1-194 of SEQ ID NO:18), 2 (amino acids 195-387 of SEQ ID NO:18), 3 (amino acids 388-585 of SEQ ID NO:18), 1 + 2 (1-387 of SEQ ID NO:18), 2 + 3 (195-585 of SEQ ID NO:18) or 1 + 3 (amino acids 1-194 of SEQ ID NO:18 + amino acids 388-585 of SEQ ID NO:18). Each

domain is itself made up of two homologous subdomains namely 1-105, 120-194, 195-291, 316-387, 388-491 and 512-585, with flexible inter-subdomain linker regions comprising residues Lys106 to Glu119, Glu292 to Va1315 and Glu492 to Ala511.

Preferably, the albumin portion of an albumin fusion protein of the invention comprises at least one subdomain or domain of HA or conservative modifications thereof. If the fusion is based on subdomains, some or all of the adjacent linker is preferably used to link to the Therapeutic protein moiety.

Albumin Fusion Proteins

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The present invention relates generally to albumin fusion proteins and methods of treating, preventing, or ameliorating diseases or disorders. As used herein, "albumin fusion protein" refers to a protein formed by the fusion of at least one molecule of albumin (or a fragment or variant thereof) to at least one molecule of a Therapeutic protein (or fragment or variant thereof). An albumin fusion protein of the invention comprises at least a fragment or variant of a Therapeutic protein and at least a fragment or variant of human serum albumin, which are associated with one another, preferably by genetic fusion (i.e., the albumin fusion protein is generated by translation of a nucleic acid in which a polynucleotide encoding all or a portion of a Therapeutic protein is joined in-frame with a polynucleotide encoding all or a portion of albumin) or chemical conjugation to one another. The Therapeutic protein and albumin protein, once part of the albumin fusion protein, may be referred to as a "portion", "region" or "moiety" of the albumin fusion protein.

In one embodiment, the invention provides an albumin fusion protein comprising, or alternatively consisting of, a Therapeutic protein (e.g., as described in Table 1) and a serum albumin protein. In other embodiments, the invention provides an albumin fusion protein comprising, or alternatively consisting of, a biologically active and/or therapeutically active fragment of a Therapeutic protein and a serum albumin protein. In other embodiments, the invention provides an albumin fusion protein comprising, or alternatively consisting of, a biologically active and/or therapeutically active variant of a Therapeutic protein and a serum albumin protein. In preferred embodiments, the serum albumin protein component of the albumin fusion protein is the mature portion of serum albumin.

In further embodiments, the invention provides an albumin fusion protein comprising, or alternatively consisting of, a Therapeutic protein, and a biologically active and/or therapeutically active fragment of serum albumin. In further embodiments, the invention provides an albumin fusion protein comprising, or alternatively consisting of, a Therapeutic protein and a biologically active and/or therapeutically active variant of serum albumin. In preferred embodiments, the Therapeutic protein portion of the albumin fusion protein is the mature portion of the Therapeutic protein.

In further embodiments, the invention provides an albumin fusion protein comprising, or alternatively consisting of, a biologically active and/or therapeutically active fragment or variant of a Therapeutic protein and a biologically active and/or therapeutically active fragment or variant of serum albumin. In preferred embodiments, the invention provides an albumin fusion protein comprising, or alternatively consisting of, the mature portion of a Therapeutic protein and the mature portion of serum albumin.

Preferably, the albumin fusion protein comprises HA as the N-terminal portion, and a Therapeutic protein as the C-terminal portion. Alternatively, an albumin fusion protein comprising HA as the C-terminal portion, and a Therapeutic protein as the N-terminal portion may also be used.

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In other embodiments, the albumin fusion protein has a Therapeutic protein fused to both the N-terminus and the C-terminus of albumin. In a preferred embodiment, the Therapeutic proteins fused at the N- and C- termini are the same Therapeutic proteins. In a preferred embodiment, the Therapeutic proteins fused at the N- and C- termini are different Therapeutic proteins fused at the N- and C- termini are different Therapeutic proteins which may be used to treat or prevent the same disease, disorder, or condition (e.g. as listed in the "Preferred Indication Y" column of Table 1). In another preferred embodiment, the Therapeutic proteins fused at the N- and C-termini are different Therapeutic proteins which may be used to treat or prevent diseases or disorders (e.g. as listed in the "Preferred Indication Y" column of Table 1) which are known in the art to commonly occur in patients simultaneously.

In addition to albumin fusion protein in which the albumin portion is fused N-terminal and/or C-terminal of the Therapeutic protein portion, albumin fusion proteins of the invention may also be produced by inserting the Therapeutic protein or peptide of interest (e.g., Therapeutic protein X as diclosed in Table 1) into an internal region of HA. For instance, within the protein sequence of the HA molecule a number of loops or turns exist between the end and beginning of α-helices, which are stabilized by disulphide bonds (see Figures 9-11). The loops, as determined from the crystal structure of HA (Fig. 13) (PDB identifiers 1AO6, 1BJ5, 1BKE, 1BMO, 1E7E to 1E7I and 1UOR) for the most part extend away from the body of the molecule. These loops are useful for the insertion, or internal fusion, of therapeutically active peptides, particularly those requiring a secondary structure to be functional, or Therapeutic proteins, to essentially generate an albumin molecule with specific biological activity.

Loops in human albumin structure into which peptides or polypeptides may be inserted to generate albumin fusion proteins of the invention include: Val54-Asn61, Thr76-Asp89, Ala92-Glu100, Gln170-Ala176, His247-Glu252, Glu266-Glu277, Glu280-His288, Ala362-Glu368, Lys439-Pro447, Val462-Lys475, Thr478-Pro486, and Lys560-Thr566. In

more preferred embodiments, peptides or polypeptides are inserted into the Val54-Asn61, Gln170-Ala176, and/or Lys560-Thr566 loops of mature human albumin (SEQ ID NO:18).

Peptides to be inserted may be derived from either phage display or synthetic peptide libraries screened for specific biological activity or from the active portions of a molecule with the desired function. Additionally, random peptide libraries may be generated within particular loops or by insertions of randomized peptides into particular loops of the HA molecule and in which all possible combinations of amino acids are represented.

Such library(s) could be generated on HA or domain fragments of HA by one of the following methods:

(a) randomized mutation of amino acids within one or more peptide loops of HA or HA domain fragments. Either one, more or all the residues within a loop could be mutated in this manner (for example see Fig. 10a);

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- (b) replacement of, or insertion into one or more loops of HA or HA domain fragments (i.e., internal fusion) of a randomized peptide(s) of length X_n (where X is an amino acid and n is the number of residues (for example see Fig. 10b);
- (c) N-, C- or N- and C- terminal peptide/protein fusions in addition to (a) and/or (b).

The HA or HA domain fragment may also be made multifunctional by grafting the peptides derived from different screens of different loops against different targets into the same HA or HA domain fragment.

In preferred embodiments, peptides inserted into a loop of human serum albumin are peptide fragments or peptide variants of the Therapeutic proteins disclosed in Table 1. More particulary, the invention encompasses albumin fusion proteins which comprise peptide fragments or peptide variants at least 7 at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 20, at least 25, at least 30, at least 35, or at least 40 amino acids in length inserted into a loop of human serum albumin. The invention also encompasses albumin fusion proteins which comprise peptide fragments or peptide variants at least 7 at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 20, at least 25, at least 30, at least 35, or at least 40 amino acids fused to the N-terminus of human serum albumin. The invention also encompasses albumin fusion proteins which comprise peptide fragments or peptide variants at least 7 at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 30, at least 35, or at least 35, or at least 35, or at least 35, or at least 40 amino acids fused to the C-terminus of human serum albumin.

Generally, the albumin fusion proteins of the invention may have one HA-derived region and one Therapeutic protein-derived region. Multiple regions of each protein, however, may be used to make an albumin fusion protein of the invention. Similarly, more than one Therapeutic protein may be used to make an albumin fusion protein of the invention.

For instance, a Therapeutic protein may be fused to both the N- and C-terminal ends of the HA. In such a configuration, the Therapeutic protein portions may be the same or different Therapeutic protein molecules. The structure of bifunctional albumin fusion proteins may be represented as: X-HA-Y or Y-HA-X.

For example, an anti-BLySTM scFv-HA-IFN α -2b fusion may be prepared to modulate the immune response to IFN α -2b by anti-BLySTM scFv. An alternative is making a bi (or even multi) functional dose of HA-fusions *e.g.* HA-IFN α -2b fusion mixed with HA-anti-BLySTM scFv fusion or other HA-fusions in various ratio's depending on function, half-life etc.

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Bi- or multi-functional albumin fusion proteins may also be prepared to target the Therapeutic protein portion of a fusion to a target organ or cell type via protein or peptide at the opposite terminus of HA.

As an alternative to the fusion of known therapeutic molecules, the peptides could be obtained by screening libraries constructed as fusions to the N-, C- or N- and C- termini of HA, or domain fragment of HA, of typically 6, 8, 12, 20 or 25 or X_n (where X is an amino acid (aa) and n equals the number of residues) randomized amino acids, and in which all possible combinations of amino acids were represented. A particular advantage of this approach is that the peptides may be selected *in situ* on the HA molecule and the properties of the peptide would therefore be as selected for rather than, potentially, modified as might be the case for a peptide derived by any other method then being attached to HA.

Additionally, the albumin fusion proteins of the invention may include a linker peptide between the fused portions to provide greater physical separation between the moieties and thus maximize the accessibility of the Therapeutic protein portion, for instance, for binding to its cognate receptor. The linker peptide may consist of amino acids such that it is flexible or more rigid.

The linker sequence may be cleavable by a protease or chemically to yield the growth hormone related moiety. Preferably, the protease is one which is produced naturally by the host, for example the S. cerevisiae protease kex2 or equivalent proteases.

Therefore, as described above, the albumin fusion proteins of the invention may have the following formula R1-L-R2; R2-L-R1; or R1-L-R2-L-R1, wherein R1 is at least one Therapeutic protein, peptide or polypeptide sequence, and not necessarily the same Therapeutic protein, L is a linker and R2 is a serum albumin sequence.

In preferred embodiments, Albumin fusion proteins of the invention comprising a Therapeutic protein have extended shelf life compared to the shelf life the same Therapeutic protein when not fused to albumin. Shelf-life typically refers to the time period over which the therapeutic activity of a Therapeutic protein in solution or in some other storage

formulation, is stable without undue loss of therapeutic activity. Many of the Therapeutic proteins are highly labile in their unfused state. As described below, the typical shelf-life of these Therapeutic proteins is markedly prolonged upon incorporation into the albumin fusion protein of the invention.

Albumin fusion proteins of the invention with "prolonged" or "extended" shelf-life exhibit greater therapeutic activity relative to a standard that has been subjected to the same storage and handling conditions. The standard may be the unfused full-length Therapeutic protein. When the Therapeutic protein portion of the albumin fusion protein is an analog, a variant, or is otherwise altered or does not include the complete sequence for that protein, the prolongation of therapeutic activity may alternatively be compared to the unfused equivalent of that analog, variant, altered peptide or incomplete sequence. As an example, an albumin fusion protein of the invention may retain greater than about 100% of the therapeutic activity, or greater than about 105%, 110%, 120%, 130%, 150% or 200% of the therapeutic activity of a standard when subjected to the same storage and handling conditions as the standard when compared at a given time point.

Shelf-life may also be assessed in terms of therapeutic activity remaining after storage, normalized to therapeutic activity when storage began. Albumin fusion proteins of the invention with prolonged or extended shelf-life as exhibited by prolonged or extended therapeutic activity may retain greater than about 50% of the therapeutic activity, about 60%, 70%, 80%, or 90% or more of the therapeutic activity of the equivalent unfused Therapeutic protein when subjected to the same conditions. For example, as discussed in Example 1, an albumin fusion protein of the invention comprising hGH fused to the full length HA sequence may retain about 80% or more of its original activity in solution for periods of up to 5 weeks or more under various temperature conditions.

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Expression of Fusion Proteins

The albumin fusion proteins of the invention may be produced as recombinant molecules by secretion from yeast, a microorganism such as a bacterium, or a human or animal cell line. Preferably, the polypeptide is secreted from the host cells. We have found that, by fusing the hGH coding sequence to the HA coding sequence, either to the 5' end or 3' end, it is possible to secrete the albumin fusion protein from yeast without the requirement for a yeast-derived pro sequence. This was surprising, as other workers have found that a yeast derived pro sequence was needed for efficient secretion of hGH in yeast.

For example, Hiramatsu *et al.* (Appl Environ Microbiol 56:2125 (1990); Appl Environ Microbiol 57:2052 (1991)) found that the N-terminal portion of the pro sequence in the *Mucor pusillus* rennin pre-pro leader was important. Other authors, using the MF α -1 signal, have always included the MF α -1 pro sequence when secreting hGH. The pro sequences were

believed to assist in the folding of the hGH by acting as an intramolecular chaperone. The present invention shows that HA or fragments of HA can perform a similar function.

Hence, a particular embodiment of the invention comprises a DNA construct encoding a signal sequence effective for directing secretion in yeast, particularly a yeast-derived signal sequence (especially one which is homologous to the yeast host), and the fused molecule of the first aspect of the invention, there being no yeast-derived pro sequence between the signal and the mature polypeptide.

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The Saccharomyces cerevisiae invertase signal is a preferred example of a yeast-derived signal sequence.

Conjugates of the kind prepared by Poznansky *et al.*, (FEBS Lett. 239:18 (1988)), in which separately-prepared polypeptides are joined by chemical cross-linking, are not contemplated.

The present invention also includes a cell, preferably a yeast cell transformed to express an albumin fusion protein of the invention. In addition to the transformed host cells themselves, the present invention also contemplates a culture of those cells, preferably a monoclonal (clonally homogeneous) culture, or a culture derived from a monoclonal culture, in a nutrient medium. If the polypeptide is secreted, the medium will contain the polypeptide, with the cells, or without the cells if they have been filtered or centrifuged away. Many expression systems are known and may be used, including bacteria (for example *E. coli* and *Bacillus subtilis*), yeasts (for example *Saccharomyces cerevisiae*, *Kluyveromyces lactis* and *Pichia pastoris*, filamentous fungi (for example *Aspergillus*), plant cells, animal cells and insect cells.

Preferred yeast strains to be used in the production of albumin fusion proteins are D88, DXY1 and BXP10. D88 [leu2-3, leu2-122, can1, pra1, ubc4] is a derivative of parent strain AH22his* (also known as DB1; see, e.g., Sleep et al. Biotechnology 8:42-46 (1990)). The strain contains a leu2 mutation which allows for auxotropic selection of 2 micron-based plasmids that contain the LEU2 gene. D88 also exhibits a derepression of PRB1 in glucose excess. The PRB1 promoter is normally controlled by two checkpoints that monitor glucose levels and growth stage. The promoter is activated in wild type yeast upon glucose depletion and entry into stationary phase. Strain D88 exhibits the repression by glucose but maintains the induction upon entry into stationary phase. The PRA1 gene encodes a yeast vacuolar protease, YscA endoprotease A, that is localized in the ER. The UBC4 gene is in the ubiquitination pathway and is involved in targeting short lived and abnormal proteins for ubiquitin dependant degradation. Isolation of this ubc4 mutation was found to increase the copy number of an expression plasmid in the cell and cause an increased level of expression of a desired protein expressed from the plasmid (see, e.g., International Publication No. WO99/00504, hereby incorporated in its entirety by reference herein).

DXY1, a derivative of D88, has the following genotype: [leu2-3, leu2-122, can1, pra1, ubc4, ura3::yap3]. In addition to the mutations isolated in D88, this strain also has a knockout of the YAP3 protease. This protease causes cleavage of mostly di-basic residues (RR, RK, KR, KK) but can also promote cleavage at single basic residues in proteins. Isolation of this yap3 mutation resulted in higher levels of full length HSA production (see, e.g., U.S. Patent No. 5,965,386, and Kerry-Williams et al., Yeast 14:161-169 (1998), hereby incorporated in their entireties by reference herein).

BXP10 has the following genotype: leu2-3, leu2-122, can1, pra1, ubc4, ura3, yap3::URA3, lys2, hsp150::LYS2, pmt1::URA3. In addition to the mutations isolated in DXY1, this strain also has a knockout of the PMT1 gene and the HSP150 gene. The PMT1 gene is a member of the evolutionarily conserved family of dolichyl-phosphate-D-mannose protein O-mannosyltransferases (Pmts). The transmembrane topology of Pmt1p suggests that it is an integral membrane protein of the endoplasmic reticulum with a role in O-linked glycosylation. This mutation serves to reduce/eliminate O-linked glycosylation of HSA fusions (see, e.g., International Publication No. WO00/44772, hereby incorporated in its entirety by reference herein). Studies revealed that the Hsp150 protein is inefficiently separated from rHA by ion exchange chromatography. The mutation in the HSP150 gene removes a potential contaminant that has proven difficult to remove by standard purification techniques. See, e.g., U.S. Patent No. 5,783,423, hereby incorporated in its entirety by reference herein.

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The desired protein is produced in conventional ways, for example from a coding sequence inserted in the host chromosome or on a free plasmid. The yeasts are transformed with a coding sequence for the desired protein in any of the usual ways, for example electroporation. Methods for transformation of yeast by electroporation are disclosed in Becker & Guarente (1990) *Methods Enzymol*. 194, 182.

Successfully transformed cells, i.e., cells that contain a DNA construct of the present invention, can be identified by well known techniques. For example, cells resulting from the introduction of an expression construct can be grown to produce the desired polypeptide. Cells can be harvested and lysed and their DNA content examined for the presence of the DNA using a method such as that described by Southern (1975) J. Mol. Biol. 98, 503 or Berent et al. (1985) Biotech. 3, 208. Alternatively, the presence of the protein in the supernatant can be detected using antibodies.

Useful yeast plasmid vectors include pRS403-406 and pRS413-416 and are generally available from Stratagene Cloning Systems, La Jolla, CA 92037, USA. Plasmids pRS403, pRS404, pRS405 and pRS406 are Yeast Integrating plasmids (YIps) and incorporate the yeast selectable markers HIS3, 7RP1, LEU2 and URA3. Plasmids pRS413-416 are Yeast Centromere plasmids (Ycps).

Preferred vectors for making albumin fusion proteins for expression in yeast include pPPC0005, pScCHSA, pScNHSA, and pC4:HSA which are described in detail in Example 2. Figure 4 shows a map of the pPPC0005 plasmid that can be used as the base vector into which polynucleotides encoding Therapeutic proteins may be cloned to form HA-fusions. It contains a PRB1 S. cerevisiae promoter (PRB1p), a Fusion leader sequence (FL), DNA encoding HA (rHA) and an ADH1 S. cerevisiae terminator sequence. The sequence of the fusion leader sequence consists of the first 19 ámino acids of the signal peptide of human serum albumin (SEQ ID NO:29) and the last five amino acids of the mating factor alpha 1 promoter (SLDKR, see EP-A-387 319 which is hereby incorporated by reference in its entirety.

The plasmids, pPPC0005, pScCHSA, pScNHSA, and pC4:HSA were deposited on April 11, 2001 at the American Type Culture Collection, 10801 University Boulevard, Manassas, Virginia 20110-2209 and given accession numbers ATCC _____, ____, and _____, respectively. Another vector useful for expressing an albumin fusion protein in yeast the pSAC35 vector which is described in Sleep *et al.*, BioTechnology 8:42 (1990) which is hereby incorporated by reference in its entirety.

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A variety of methods have been developed to operably link DNA to vectors via complementary cohesive termini. For instance, complementary homopolymer tracts can be added to the DNA segment to be inserted to the vector DNA. The vector and DNA segment are then joined by hydrogen bonding between the complementary homopolymeric tails to form recombinant DNA molecules.

Synthetic linkers containing one or more restriction sites provide an alternative method of joining the DNA segment to vectors. The DNA segment, generated by endonuclease restriction digestion, is treated with bacteriophage T4 DNA polymerase or E. coli DNA polymerase I, enzymes that remove protruding, γ-single-stranded termini with their 3' 5'-exonucleolytic activities, and fill in recessed 3'-ends with their polymerizing activities.

The combination of these activities therefore generates blunt-ended DNA segments. The blunt-ended segments are then incubated with a large molar excess of linker molecules in the presence of an enzyme that is able to catalyze the ligation of blunt-ended DNA molecules, such as bacteriophage T4 DNA ligase. Thus, the products of the reaction are DNA segments carrying polymeric linker sequences at their ends. These DNA segments are then cleaved with the appropriate restriction enzyme and ligated to an expression vector that has been cleaved with an enzyme that produces termini compatible with those of the DNA segment.

Synthetic linkers containing a variety of restriction endonuclease sites are commercially available from a number of sources including International Biotechnologies Inc, New Haven, CT, USA.

A desirable way to modify the DNA in accordance with the invention, if, for example,

HA variants are to be prepared, is to use the polymerase chain reaction as disclosed by Saiki et al. (1988) Science 239, 487-491. In this method the DNA to be enzymatically amplified is flanked by two specific oligonucleotide primers which themselves become incorporated into the amplified DNA. The specific primers may contain restriction endonuclease recognition sites which can be used for cloning into expression vectors using methods known in the art.

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Exemplary genera of yeast contemplated to be useful in the practice of the present invention as hosts for expressing the albumin fusion proteins are Pichia (formerly classified as Hansenula), Saccharomyces, Kluyveromyces, Aspergillus, Candida, Torulopsis, Citeromyces, Pachysolen, Zygosaccharomyces, Schizosaccharomyces, Torulaspora, Debaromyces, Trichoderma, Cephalosporium, Humicola, Mucor, Neurospora, Yarrowia. Leucosporidium, Botryoascus, Sporidiobolus. Rhodosporidium, Metschunikowia, Endomycopsis, and the like. Preferred genera are those selected from the group consisting of Saccharomyces, Schizosaccharomyces, Kluyveromyces, Pichia and Torulaspora. Examples of Saccharomyces spp. are S. cerevisiae, S. italicus and S. rouxii.

Examples of Kluyveromyces spp. are K. fragilis, K. lactis and K. marxianus. A suitable Torulaspora species is T. delbrueckii. Examples of Pichia (Hansenula) spp. are P. angusta (formerly H. polymorpha), P. anomala (formerly H. anomala) and P. pastoris. Methods for the transformation of S. cerevisiae are taught generally in EP 251 744, EP 258 067 and WO 90/01063, all of which are incorporated herein by reference.

Preferred exemplary species of Saccharomyces include S. cerevisiae, S. italicus, S. diastaticus, and Zygosaccharomyces rouxii. Preferred exemplary species of Kluyveromyces include K. fragilis and K. lactis. Preferred exemplary species of Hansenula include H. polymorpha (now Pichia angusta), H. anomala (now Pichia anomala), and Pichia capsulata. Additional preferred exemplary species of Pichia include P. pastoris. Preferred exemplary species of Aspergillus include A. niger and A. nidulans. Preferred exemplary species of Yarrowia include Y. lipolytica. Many preferred yeast species are available from the ATCC. For example, the following preferred yeast species are available from the ATCC and are useful in the expression of albumin fusion proteins: Saccharomyces cerevisiae Hansen, teleomorph strain BY4743 yap3 mutant (ATCC Accession No. 4022731); Saccharomyces cerevisiae Hansen, teleomorph strain BY4743 hsp150 mutant (ATCC Accession No. 4021266); Saccharomyces cerevisiae Hansen, teleomorph strain BY4743 pmt1 mutant (ATCC Accession No. 4023792); Saccharomyces cerevisiae Hansen, teleomorph (ATCC Accession Nos. 20626; 44773; 44774; and 62995); Saccharomyces diastaticus Andrews et Gilliland ex van der Walt, teleomorph (ATCC Accession No. 62987); Kluyveromyces lactis (Dombrowski) van der Walt, teleomorph (ATCC Accession No. 76492); Pichia angusta (Teunisson et al.) Kurtzman, teleomorph deposited as Hansenula polymorpha de Morais et Maia, teleomorph (ATCC Accession No. 26012); Aspergillus niger van Tieghem, anamorph

(ATCC Accession No. 9029); Aspergillus niger van Tieghem, anamorph (ATCC Accession No. 16404); Aspergillus nidulans (Eidam) Winter, anamorph (ATCC Accession No. 48756); and Yarrowia lipolytica (Wickerham et al.) van der Walt et von Arx, teleomorph (ATCC Accession No. 201847).

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Suitable promoters for S. cerevisiae include those associated with the PGKI gene, GAL1 or GAL10 genes, CYCI, PHO5, TRPI, ADHI, ADH2, the genes for glyceraldehyde-3-phosphate dehydrogenase, hexokinase, pyruvate decarboxylase, phosphofructokinase, triose phosphate isomerase, phosphoglucose isomerase, glucokinase, alpha-mating factor pheromone, [a mating factor pheromone], the PRBI promoter, the GUT2 promoter, the GPDI promoter, and hybrid promoters involving hybrids of parts of 5' regulatory regions with parts of 5' regulatory regions of other promoters or with upstream activation sites (e.g. the promoter of EP-A-258 067).

Convenient regulatable promoters for use in *Schizosaccharomyces pombe* are the thiamine-repressible promoter from the nmt gene as described by Maundrell (1990) *J. Biol. Chem.* 265, 10857-10864 and the glucose repressible jbpl gene promoter as described by Hoffman & Winston (1990) *Genetics* 124, 807-816.

Methods of transforming *Pichia* for expression of foreign genes are taught in, for example, Cregg *et al.* (1993), and various Phillips patents (*e.g.* US 4 857 467, incorporated herein by reference), and *Pichia* expression kits are commercially available from Invitrogen BV, Leek, Netherlands, and Invitrogen Corp., San Diego, California. Suitable promoters include AOXI and AOX2. Gleeson *et al.* (1986) J. Gen. Microbiol. 132, 3459-3465 include information on *Hansenula* vectors and transformation, suitable promoters being MOX1 and FMD1; whilst EP 361 991, Fleer *et al.* (1991) and other-publications from Rhone-Poulenc Rorer teach how to express foreign proteins in *Kluyveromyces* spp., a suitable promoter being PGKI.

The transcription termination signal is preferably the 3' flanking sequence of a eukaryotic gene which contains proper signals for transcription termination and polyadenylation. Suitable 3' flanking sequences may, for example, be those of the gene naturally linked to the expression control sequence used, *i.e.* may correspond to the promoter. Alternatively, they may be different in which case the termination signal of the S. cerevisiae ADHI gene is preferred.

The desired albumin fusion protein may be initially expressed with a secretion leader sequence, which may be any leader effective in the yeast chosen. Leaders useful in S. cerevisiae include that from the mating factor α polypeptide (MF α -1) and the hybrid leaders of EP-A-387 319. Such leaders (or signals) are cleaved by the yeast before the mature albumin is released into the surrounding medium. Further such leaders include those of S. cerevisiae invertase (SUC2) disclosed in JP 62-096086 (granted as 911036516), acid

phosphatase (PH05), the pre-sequence of MFα-1, 0 glucanase (BGL2) and killer toxin; S. diastaticus glucoarnylase II; S. carlsbergensis α-galactosidase (MEL1); K. lactis killer toxin; and Candida glucoarnylase.

Additional Methods of Recombinant and Synthetic Production of Albumin Fusion Proteins

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The present invention also relates to vectors containing a polynucleotide encoding an albumin fusion protein of the present invention, host cells, and the production of albumin fusion proteins by synthetic and recombinant techniques. The vector may be, for example, a phage, plasmid, viral, or retroviral vector. Retroviral vectors may be replication competent or replication defective. In the latter case, viral propagation generally will occur only in complementing host cells.

The polynucleotides encoding albumin fusion proteins of the invention may be joined to a vector containing a selectable marker for propagation in a host. Generally, a plasmid vector is introduced in a precipitate, such as a calcium phosphate precipitate, or in a complex with a charged lipid. If the vector is a virus, it may be packaged in vitro using an appropriate packaging cell line and then transduced into host cells.

The polynucleotide insert should be operatively linked to an appropriate promoter, such as the phage lambda PL promoter, the E. coli lac, trp, phoA and tac promoters, the SV40 early and late promoters and promoters of retroviral LTRs, to name a few. Other suitable promoters will be known to the skilled artisan. The expression constructs will further contain sites for transcription initiation, termination, and, in the transcribed region, a ribosome binding site for translation. The coding portion of the transcripts expressed by the constructs will preferably include a translation initiating codon at the beginning and a termination codon (UAA, UGA or UAG) appropriately positioned at the end of the polypeptide to be translated.

As indicated, the expression vectors will preferably include at least one selectable marker. Such markers include dihydrofolate reductase, G418, glutamine synthase, or neomycin resistance for eukaryotic cell culture, and tetracycline, kanamycin or ampicillin resistance genes for culturing in *E. coli* and other bacteria. Representative examples of appropriate hosts include, but are not limited to, bacterial cells, such as *E. coli*, Streptomyces and Salmonella typhimurium cells; fungal cells, such as yeast cells (e.g., Saccharomyces cerevisiae or Pichia pastoris (ATCC Accession No. 201178)); insect cells such as Drosophila S2 and Spodoptera Sf9 cells; animal cells such as CHO, COS,NSO, 293, and Bowes melanoma cells; and plant cells. Appropriate culture mediums and conditions for the above-described host cells are known in the art.

Among vectors preferred for use in bacteria include pQE70, pQE60 and pQE-9,